WORLD INTELLECTUAL PROPERTY ORGANIZATION --International Bureau



(51) International Patent Classification 6:		(11) International Publication Number: WO 97/42286
C11D 3/37	A1	(43) International Publication Date: 13 November 1997 (13.11.97)
(21) International Application Number: PCT/US (22) International Filing Date: 25 April 1997 ((AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU,
(30) Priority Data: 60/016,525 3 May 1996 (03.05.96)		Published With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of
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LIQUID LAUNDRY DETERGENT COMPOSITIONS COMPRISING COTTON SOIL RELEASE POLYMERS

FIELD OF THE INVENTION

The present invention relates to liquid laundry detergent compositions comprising water soluble and/or dispersible, modified polyamines having functionalized backbone moieties which provide cotton soil release benefits in combination with selected non-cotton soil release agents.

BACKGROUND OF THE INVENTION

A wide variety of soil release agents for use in domestic and industrial fabric treatment processes such as laundering, fabric drying in hot air clothes dryers, and the like are known in the art. Various soil release agents have been commercialized and are currently used in detergent compositions and fabric softener/antistatic articles and compositions. Such soil release polymers typically comprise an oligomeric or polymeric ester "backbone".

Soil release polymers are generally very effective on polyester or other synthetic fabrics where the grease, oil or similar hydrophobic stains spread out and form a attached film and thereby are not easily removed in an aqueous laundering process. Many soil release polymers have a less dramatic effect on "blended" fabrics, that is on fabrics that comprise a mixture of cotton and synthetic material, and have little or no effect on cotton articles. The reason for the affinity of many soil release agents for synthetic fabric is that the backbone of a polyester soil release polymer typically comprises a mixture of terephthalate residues and ethyleneoxy or propyleneoxy polymeric units; the same or closely analogous to materials that comprise the polyester fibers of synthetic fabric. This similar structure of soil release agents and synthetic fabric produce an intrinsic affinity between these compounds.

Extensive research in this area has yielded significant improvements in the effectiveness of polyester soil release agents yielding materials with enhanced product performance and formulatability. Modifications of the polymer backbone as well as the selection of proper end-capping groups has produced a wide variety of polyester soil release polymers. For example, end-cap modifications, such as the use of sulfoaryl moieties and especially the low cost isethionate-derived end-capping units, have increased the range of solubility and adjunct ingredient compatibility of these polymers without sacrifice of soil release effectiveness. Many polyester soil release polymers can now be formulated into both liquid as well as solid (i.e., granular) detergents.

In contrast to the case of polyester soil release agents, producing an oligomeric or polymeric material that mimics the structure of cotton has not resulted in a cotton soil release polymer. Although cotton and polyester fabric are both comprised of long chain polymeric materials, they are chemically very different. Cotton is comprised of cellulose fibers that consist of anhydroglucose units joined by 1-4 linkages. These glycosidic linkages characterize the cotton cellulose as a polysaccharide whereas polyester soil release polymers are generally a combination of terephthalate and oxyethylene/oxypropylene residues. These differences in composition account for the difference in the fabric properties of cotton versus polyester fabric. Cotton is hydrophilic relative to polyester. Polyester is hydrophobic and attracts oily or greasy dirt and can easily be "dry cleaned". Importantly, the terephthalate and ethyleneoxy/propyleneoxy backbone of polyester fabric does not contain reactive sites, such as the hydroxyl moieties of cotton, that interact with stains in a different manner than synthetics. Many cotton stains become "fixed" and can only be resolved by bleaching the fabric.

Until now the development of an effective cotton soil release agent for use in a laundry detergent has been elusive. Attempts by others to apply the paradigm of matching the structure of a soil release polymer with the structure of the fabric, a method successful in the polyester soil release polymer field, has nevertheless yielded marginal results when applied to cotton fabric soil release agents. The use of methylcellulose, a cotton polysaccharide with modified oligomeric units, proved to be more effective on polyesters than on cotton.

For example, U.K. 1,314,897, published April 26, 1973 teaches a hydroxypropyl methyl cellulose material for the prevention of wet-soil redeposition and improving stain release on laundered fabric. While this material appears to be somewhat effective on polyester and blended fabrics, the disclosure indicates these materials to be unsatisfactory at producing the desired results on cotton fabric.

Other attempts to produce a soil release agent for cotton fabric have usually taken the form of permanently modifying the chemical structure of the cotton fibers themselves by reacting a substrate with the polysaccharide polymer backbone. For example, U. S. Patent No. 3,897,026 issued to Kearney, discloses cellulosic textile materials having improved soil release and stain resistance properties obtained by reaction of an ethylenemaleic anhydride co-polymer with the hydroxyl moieties of the cotton polymers. One perceived drawback of this method is the desirable hydrophilic properties of the cotton fabric are substantially modified by this process.

Non-permanent soil release treatments or finishes have also been previously attempted. U.S. Patent No. 3,912,681 issued to Dickson teaches a composition for

applying a non-permanent soil release finish comprising a polycarboxylate polymer to a cotton fabric. However, this material must be applied at a pH less than 3, a process not suitable for consumer use nor compatible with laundry detergents which typically have a pH greater than 7.5.

U.S. Patent No. 3,948,838 issued to Hinton, *et alia* describes high molecular weight (500,000 to 1,500,000) polyacrylic polymers for soil release. These materials are used preferably with other fabric treatments, for example, durable press textile reactants such as formaldehyde. This process is also not readily applicable for use by consumers in a typical washing machine.

U.S. Patent 4,559,056 issued to Leigh, et alia discloses a process for treating cotton or synthetic fabrics with a composition comprising an organopolysiloxane elastomer, an organosiloxaneoxyalkylene copolymer crosslinking agent and a siloxane curing catalyst. Organosilicone oligomers are well known by those skilled in the art as suds supressors

Other soil release agents not comprising terephthalate and mixtures of polyoxy ethylene/propylene are vinyl caprolactam resins as disclosed by Rupert, et alia in U.S. Patent Nos. 4,579,681 and 4,614,519. These disclosed vinyl caprolactam materials have their effectiveness limited to polyester fabrics, blends of cotton and polyester, and cotton fabrics rendered hydrophobic by finishing agents.

Examples of alkoxylated polyamines and quaternized alkoxylated polyamines are disclosed in European Patent Application 206,513 as being suitable for use as soil dispersents, however their possible use as a cotton soil release agent is not disclosed. In addition, these materials do not comprise N-oxides, a key modification made to the polyamines of the present invention and a component of the increased bleach stability exhibited by the presently disclosed compounds.

It has now been surprisingly discovered that effective soil release agents for cotton articles can be prepared from certain modified polyamines. This unexpected result has yielded compositions that are effective at providing the soil release benefits once available to only synthetic and synthetic-cotton blended fabric. When the cotton soil release polymers of the present invention are used in combination with non-cotton soil release agents, the full spectrum of fabric types is provided with soil release benefits.

The present invention provides for liquid laundry detergent compositions that comprise nonionic and anionic surfactants together with a combination of non-cotton soil release polymers and the cotton soil release agents of the present invention. These combinations provide a liquid laundry detergent composition that is effective for providing soil release benefits to all fabric. The liquid detergents can have a wide range

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of viscosity and may include heavy concentrates, pourable "ready" detergents, or light duty fabric pre-treatments.

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BACKGROUND ART

In addition to the above cited art, the following disclose various soil release polymers or modified polyamines; U.S. Patent 4,548,744, Connor, issued October 22, 1985; U.S. Patent 4,597,898, Vander Meer, issued July 1, 1986; U.S. Patent 4,877,896, Maldonado, et al., issued October 31, 1989; U.S. Patent 4,891,160, Vander Meer, issued January 2, 1990; U.S. Patent 4,976,879, Maldonado, et al., issued December 11, 1990; U.S. Patent 5,415,807, Gosselink, issued May 16,1995; U.S. Patent 4,235,735, Marco, et al., issued November 25, 1980; U.K. Patent 1,537,288, published December 29, 1978; U.K. Patent 1,498,520, published January 18, 1978; WO 95/32272, published November 30, 1995; German Patent DE 28 29 022, issued January 10, 1980; Japanese Kokai JP 06313271, published April 27, 1994.

SUMMARY OF THE INVENTION

The present invention relates to liquid laundry detergent compositions which provide cotton soil release benefits, comprising:

- A) at least about 0.01% by weight, of an anionic detersive surfactant selected from the group consisting of alkyl sulfates, alkyl alkoxylated sulfates, and mixtures thereof;
- B) at least about 0.01% by weight, of a non-cotton soil release agent selected from the group consisting of a terephthalate co-polymer comprising:
 - i) a backbone comprising:
 - a) at least one moiety having the formula:

b) at least one moiety having the formula:

$$R^{10}$$
 R^{10} R^{10} R^{10} R^{10} R^{10} R^{10} R^{10} R^{10} R^{10}

wherein R⁹ is C₂-C₆ linear alkylene, C₃-C₆ branched alkylene, C₅-C₇ cyclic alkylene, and mixtures thereof; R¹⁰ is independently selected from hydrogen or -L-SO₃-

M⁺; wherein L is a side chain moiety selected from the group consisting of alkylene, oxyalkylene, alkyleneoxyalkylene, arylene, oxyarylene, alkyleneoxyarylene, poly(oxyalkylene), oxyalkyleneoxyarylene, poly(oxyalkylene)oxyarlyene, alkylene-poly(oxyalkylene), and mixtures thereof; M is hydrogen or a salt forming cation; i has the value of 0 or 1;

- c) at least one trifunctional, ester-forming, branching moiety;
- d) at least one 1,2-oxyalkyleneoxy moiety; and
- ii) one or more capping units comprising:
 - ethoxylated or propoxylated hydroxyethanesulfonate or ethoxylated or propoxylated hydroxypropanesulfonate units of the formula (MO₃S)(CH₂)_m(R¹¹O)_n-, where M is a salt forming cation, R¹¹ is ethylene, propylene, and mixtures thereof, m is 0 or 1, and n is from 1 to 20;
 - b) sulfoaroyl units of the formula $-(O)C(C_6H_4)(SO_3-M^+)$, wherein M is a salt forming cation;
 - c) modified poly(oxyethylene)oxy monoalkyl ether units of the formula R¹²O(CH₂CH₂O)_k-, wherein R¹² contains from 1 to 4 carbon atoms and k is from about 3 to about 100; and
 - d) ethoxylated or propoxylated phenolsulfonate end-capping units of the formula MO₃S(C₆H₄)(OR¹³)_nO-, wherein n is from 1 to 20; M is a salt-forming cation; and R¹³ is ethylene, propylene, and mixtures thereof;

a sulfonated oligomeric ester composition comprising the sulfonated product of a pre-formed, substantially linear ester oligomer, said linear ester oligomer comprising, per mole,

- i) 2 moles of terminal units wherein from about 1 mole to about 2 moles of said terminal units are derived from an olefinically unsaturated component selected from the group consisting of allyl alcohol and methallyl alcohol, and any remaining of said terminal units are other units of said linear ester oligomer;
- ii) from about 1 mole to about 4 moles of nonionic hydrophile units, said hydrophile units being derived from alkyleneoxides, said

- alkylene oxides comprising from about 50% to 100% ethylene oxide;
- from about 1.1 moles to about 20 moles of repeat units derived from an aryldicarbonyl component wherein said aryldicarbonyl component is comprised of from about 50% to 100% dimethylterephthalate, whereby the repeat units derived from said dimethylterephthalate are terephthaloyl; and
- iv) from about 0.1 moles to about 19 moles of repeat units derived from a diol component selected from the group consisting of C₂-C₄ glycols;

wherein the extent of sulfonation of said sulfonated oligomeric ester composition is such that said terminal units are chemically modified by

v) from about 1 mole to about 4 moles of terminal unit substituent groups of formula -SO_XM wherein x is 2 or 3, said terminal unit substituent groups being derived from a bisulfite component selected from the group consisting of HSO₃M wherein M is a conventional water-soluble cation;

a capped terephalate co-polymer having the formula

$$X[(OCH_2CH_2)_n(OR_5)_m][(A-R^1-A-R^2)_n(A-R^3-A-R^2)_v]$$

$$A-R^4-A[(R^5O)_m(CH_2CH_2O)_n]X$$

wherein each of the A moieties is selected from the group consisting of

and combinations thereof, each of the R^1 moieties is selected from the group consisting of 1,4-phenylene and combinations thereof with 1,3-phenylene, 1,2 phenylene, 1,8-naphthylene, 1,4-naphthylene, 2,2'-biphenylene, 4,4'-biphenylene, C_1 - C_8 alkylene, C_1 - C_8 alkenylene and mixtures thereof the R^2 moieties are each selected from the group consisting of ethylene moieties, substituted ethylene moieties having C_1 - C_4 alkyl, alkoxy substituents, and mixtures thereof; the R^3 moieties are substituted C_2 - C_{18} hydrocarbylene moieties having at least one - CO_2M , - $O[(R^5O)_m(CH_2CH_2O)_n]X$ or - $A[(R^2-A-R^4-A)]_w[(R^5O)_m(CH_2CH_2O)_n]X$ substituent; the R^4 moieties are R^1 or R^3 moieties, or mixtures thereof; each R^5 is C_1 - C_4 alkylene, or the moiety -

C)

 R^2 -A-R⁶- wherein R⁶ is a C_1 - C_{12} alkylene, alkenylene, arylene, or alkarylene moiety; each M is hydrogen or a water-soluble cation; each X is C_1 - C_4 alkyl; the indices m and n have the values such that the moiety - (CH_2CH_2O) - comprises at least about 50% by weight of the moiety $[(R^5O)_m(CH_2CH_2O)_n]$, provided that when R⁵ is the moiety -R²-A-R⁶-, m is 1; each n is at least about 10; the indices u and v have the value such that the sum of u + v is from about 3 to about 25; the index w is 0 or at least 1; and when w is at least 1 u, v and w have the value such that the sum of u + v + w is from about 3 to about 25; and mixtures thereof; at least about 0.01% by weight, of a water-soluble or dispersible, modified polyamine cotton soil release agent comprising a polyamine backbone corresponding to the formula:

$$[H_2N-R]_{n+1}-[N-R]_m-[N-R]_n-NH_2$$

having a modified polyamine formula $V_{(n+1)}W_mY_nZ$ or a polyamine backbone corresponding to the formula:

$$H$$
 $[H_2N-R]_{m-k+1}$
 $[N-R]_m$
 $[N-R]_m$
 $[N-R]_k$
 $[N-R]_k$

having a modified polyamine formula $V_{(n-k+1)}W_mY_nY_kZ$, wherein k is less than or equal to n, said polyamine backbone prior to modification has a molecular weight greater than about 200 daltons, wherein

i) V units are terminal units having the formula:

ii) W units are backbone units having the formula:

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iii) Y units are branching units having the formula:

$$-N-R- \qquad \text{or} \qquad -N+R- \qquad \text{or} \qquad -N-R-$$

iv) Z units are terminal units having the formula:

wherein backbone linking R units are selected from the group consisting of C2-C12 alkylene, C4-C12 alkenylene, C3-C12 hydroxyalkylene, C4-C₁₂ dihydroxy-alkylene, C₈-C₁₂ dialkylarylene, -(R¹O)_xR¹-, - $(R^{1}O)_{x}R^{5}(OR^{1})_{x^{-}}$, - $(CH_{2}CH(OR^{2})CH_{2}O)_{z^{-}}$ $(R^{1}O)_{v}R^{1}(OCH_{2}CH(OR^{2})CH_{2})_{w}$, $-C(O)(R^{4})_{r}C(O)$ -, CH₂CH(OR²)CH₂-, and mixtures thereof; wherein R¹ is C₂-C₃ alkylene and mixtures thereof; R² is hydrogen, -(R¹O)_xB, and mixtures thereof; R³ is C₁-C₁₈ alkyl, C₇-C₁₂ arylalkyl, C₇-C₁₂ alkyl substituted aryl, C₆-C₁₂ aryl, and mixtures thereof; R⁴ is C₁-C₁₂ alkylene, C₄-C₁₂ alkenylene, C₈-C₁₂ arylalkylene, C₆-C₁₀ arylene, and mixtures thereof; R⁵ is C₁-C₁₂ alkylene, C₃-C₁₂ hydroxy-alkylene, C₄-C₁₂ dihydroxyalkylene, Cg-C12 dialkylarylene, -C(O)-, - $C(O)NHR^6NHC(O)-, -R^1(OR^1)-, -C(O)(R^4)_rC(O)-,$ CH2CH(OH)CH2-, -CH2CH(OH)CH2O(R1O)vR1-OCH2CH(OH)CH2-, and mixtures thereof; R⁶ is C₂-C₁₂ alkylene or C₆-C₁₂ arylene; E units are selected from the group consisting of hydrogen, C1-C22 alkyl, C3-C₂₂ alkenyl, C₇-C₂₂ arylalkyl, C₂-C₂₂ hydroxyalkyl, -(CH₂)_pCO₂M, - $(CH_2)_0SO_3M$, $-CH(CH_2CO_2M)-CO_2M$, $-(CH_2)_pPO_3M$, $-(R^1O)_xB$, $-(R^1O)_xB$ C(O)R³, and mixtures thereof; provided that when any E unit of a nitrogen is a hydrogen, said nitrogen is not also an N-oxide; B is hydrogen, C₁-C₆ alkyl, -(CH₂)_q-SO₃M, -(CH₂)_pCO₂M, -

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(CH₂)_q(CHSO₃M)CH₂SO₃M, -(CH₂)_q-(CHSO₂M)CH₂SO₃M, -(CH₂)_pPO₃M, -PO₃M, and mixtures thereof; M is hydrogen or a water soluble cation in sufficient amount to satisfy charge balance; X is a water soluble anion; m has the value from 4 to about 400; n has the value from 0 to about 200; p has the value from 1 to 6, q has the value from 0 to 6; r has the value of 0 or 1; w has the value 0 or 1; x has the value from 1 to 100; y has the value from 0 to 100; z has the value 0 or 1; and

D) the balance carrier and adjunct ingredients wherein said composition has a pH of about 7.2 to about 8.9 when measured as a 10% solution in water.

All percentages, ratios and proportions herein are by weight, unless otherwise specified. All temperatures are in degrees Celsius (OC) unless otherwise specified. All documents cited are in relevant part, incorporated herein by reference.

DETAILED DESCRIPTION OF THE INVENTION

The present invention comprises liquid laundry detergent compositions suitable for use with cotton, non-cotton, or mixtures of cotton and non-cotton fabric. The liquid laundry detergent compositions may optionally comprise bleaching materials.

The preferred liquid laundry detergent compositions of the present invention comprise certain anionic surfactants (preferably in combination with nonionic surfactants), and selected non-cotton soil release agents that when used in combination with the cotton soil release polymers of the present invention, provides improved cleaning and soil release benefits for all fabric. The preferred liquid laundry detergent compositions of the present invention comprise the following ingredients.

Anionic Detersive Surfactants

The compositions of the present invention comprise at least about 0.01%, preferably at least 0.1%, more preferably from about 1% to about 95%, most preferably from about 1% to about 80% by weight, of an anionic detersive surfactant selected from the group consisting of alkyl sulfates, alkyl alkoxylated sulfates, and mixtures thereof. Alkyl sulfate surfactants, either primary or secondary, are a type of anionic surfactant of importance for use herein. Alkyl sulfates have the general formula ROSO₃M wherein R preferably is a C₁₀-C₂₄ hydrocarbyl, preferably an alkyl straight or branched chain or hydroxyalkyl having a C₁₀-C₂₀ alkyl component, more preferably a C₁₂-C₁₈ alkyl or hydroxyalkyl, and M is hydrogen or a water soluble cation, e.g., an alkali metal cation (e.g., sodium potassium, lithium), substituted or unsubstituted ammonium cations such as methyl-, dimethyl-, and trimethyl ammonium and quaternary ammonium cations, e.g., tetramethyl-ammonium and dimethyl piperdinium, and cations derived from alkanolamines such as ethanolamine, diethanolamine, triethanolamine, and mixtures thereof, and the like.

Typically, alkyl chains of C₁₂-C₁₆ are preferred for lower wash temperatures (e.g., below about 50°C) and C₁₆-C₁₈ alkyl chains are preferred for higher wash temperatures (e.g., about 50°C).

Alkyl alkoxylated sulfate surfactants are another category of preferred anionic surfactant. These surfactants are water soluble salts or acids typically of the formula RO(A)_mSO₃M wherein R is an unsubstituted C₁₀-C₂₄ alkyl or hydroxyalkyl group having a C₁₀-C₂₄ alkyl component, preferably a C₁₂-C₂₀ alkyl or hydroxyalkyl, more preferably C₁₂-C₁₈ alkyl or hydroxyalkyl, A is an ethoxy or propoxy unit, m is greater than zero, typically between about 0.5 and about 6, more preferably between about 0.5 and about 3, and M is hydrogen or a water soluble cation which can be, for example, a metal cation (e.g., sodium, potassium, lithium, calcium, magnesium, etc.), ammonium or substituted-ammonium cation. Alkyl ethoxylated sulfates as well as alkyl propoxylated sulfates are contemplated herein. Specific examples of substituted ammonium cations include methyl-, dimethyl-, trimethyl-ammonium and quaternary ammonium cations, such as tetramethyl-ammonium, dimethyl piperdinium and cations derived from alkanolamines, e.g., monoethanolamine, diethanolamine, and triethanolamine, and mixtures thereof. Exemplary surfactants are C₁₂C₁₈ alkyl polyethoxylate (1.0) sulfate, C₁₂-C₁₈ alkyl polyethoxylate (2.25) sulfate, C₁₂-C₁₈ alkyl polyethoxylate (3.0) sulfate, and C₁₂-C₁₈ alkyl polyethoxylate (4.0) sulfate wherein M is conveniently selected from sodium and potassium.

Nonionic Detersive Surfactants

The compositions of the present invention preferably also comprise at least about 0.01%, preferably at least 0.1%, more preferably from about 1% to about 95%, most preferably from about 1% to about 80% by weight, of an nonionic detersive surfactant. Preferred nonionic surfactants such as C₁₂-C₁₈ alkyl ethoxylates ("AE") including the so-called narrow peaked alkyl ethoxylates and C₆-C₁₂ alkyl phenol alkoxylates (especially ethoxylates and mixed ethoxy/propoxy), block alkylene oxide condensate of C₆ to C₁₂ alkyl phenols, alkylene oxide condensates of C₈-C₂₂ alkanols and ethylene oxide/propylene oxide block polymers (PluronicTM-BASF Corp.), as well as semi polar nonionics (e.g., amine oxides and phosphine oxides) can be used in the present compositions. An extensive disclosure of these types of surfactants is found in U.S. Pat. 3,929,678, Laughlin et al., issued December 30, 1975, incorporated herein by reference.

Alkylpolysaccharides such as disclosed in U.S. Pat. 4,565,647 Llenado (incorporated herein by reference) are also preferred nonionic surfactants in the compositions of the invention.

Further preferred nonionic surfactants are the polyhydroxy fatty acid amides having the formula:

wherein R7 is C5-C31 alkyl, preferably straight chain C7-C19 alkyl or alkenyl, more preferably straight chain Co-C17 alkyl or alkenyl, most preferably straight chain C11-C15 alkyl or alkenyl, or mixtures thereof; R8 is selected from the group consisting of hydrogen, C₁-C₄ alkyl, C₁-C₄ hydroxyalkyl, preferably methyl or ethyl, more preferably methyl. Q is a polyhydroxyalkyl moiety having a linear alkyl chain with at least 3 hydroxyls directly connected to the chain, or an alkoxylated derivative thereof; preferred alkoxy is ethoxy or propoxy, and mixtures thereof. Preferred Q is derived from a reducing sugar in a reductive amination reaction. More preferably Q is a glycityl mojety. Suitable reducing sugars include glucose, fructose, maltose, lactose, galactose, mannose, and xylose. As raw materials, high dextrose corn syrup, high fructose corn syrup, and high maltose corn syrup can be utilized as well as the individual sugars listed above. These corn syrups may yield a mix of sugar components for Q. It should be understood that it is by no means intended to exclude other suitable raw materials. Q is more preferably selected from the group consisting of -CH₂(CHOH)_nCH₂OH,-CH(CH₂OH)(CHOH)_{n-1}CH₂OH, -CH₂(CHOH)₂-(CHOR')(CHOH)CH2OH, and alkoxylated derivatives thereof, wherein n is an integer from 3 to 5, inclusive, and R' is hydrogen or a cyclic or aliphatic monosaccharide. Most preferred substituents for the Q moiety are glycityls wherein n is 4, particularly -CH₂(CHOH)₄CH₂OH.

R⁷CO-N < can be, for example, cocamide, stearamide, oleamide, lauramide, myristamide, capricamide, palmitamide, tallowamide, etc.

R⁸ can be, for example, methyl, ethyl, propyl, isopropyl, butyl, 2-hydroxy ethyl, or 2-hydroxy propyl.

Q can be 1-deoxyglucityl, 2-deoxyfructityl, 1-deoxymaltityl, 1-deoxylactityl, 1-deoxymaltotriotityl, etc.

A particularly desirable surfactant of this type for use in the compositions herein is alkyl-N-methyl glucomide, a compound of the above formula wherein \mathbb{R}^7 is alkyl (preferably \mathbb{C}_{11} - \mathbb{C}_{13}), \mathbb{R}^8 , is methyl and Q is 1-deoxyglucityl.

Other sugar-derived surfactants include the N-alkoxy polyhydroxy fatty acid amides, such as C_{10} - C_{18} N-(3-methoxypropyl) glucamide. The N-propyl through N-hexyl C_{12} - C_{18} glucamides can be used for low sudsing. C_{10} - C_{20} conventional soaps may also be used. If high sudsing is desired, the branched-chain C_{10} - C_{16} soaps may be used. Other conventional useful surfactants are listed in standard texts.

For the purposes of the present invention other detersive surfactants, described herein below, may be used in the liquid laundry detergent compositions.

Non-cotton Soil Release Polymers

The non-cotton soil release polymers to be used in the laundry detergent compositions of the present invention are the following.

<u>Preferred non-cotton soil release agent - A.</u> Suitable for use in the laundry detergent compositions of the present invention are preferred non-cotton soil release polymers comprising:

- a) a backbone comprising:
 - i) at least one moiety having the formula:

ii) at least one moiety having the formula:

wherein R⁹ is C₂-C₆ linear alkylene, C₃-C₆ branched alkylene, C₅-C₇ cyclic alkylene, and mixtures thereof; R¹⁰ is independently selected from hydrogen or -L-SO₃-M⁺; wherein L is a side chain moiety selected from the group consisting of alkylene, oxyalkylene, alkyleneoxyalkylene, arylene, oxyarylene, alkyleneoxyarylene, poly(oxyalkylene), oxyalkyleneoxyarylene, poly(oxyalkylene)oxyarlyene, alkylenepoly(oxyalkylene), and mixtures thereof; M is hydrogen or a salt forming cation; i has the value of 0 or 1;

- iii) at least one trifunctional, ester-forming, branching moiety;
- iv) at least one 1,2-oxyalkyleneoxy moiety; and
- b) one or more capping units comprising:
 - i) ethoxylated or propoxylated hydroxyethanesulfonate or ethoxylated or propoxylated hydroxypropanesulfonate units of the formula (MO₃S)(CH₂)_m(R¹¹O)_n-, where M is a salt forming

cation such as sodium or tetralkylammonium, R¹¹ is ethylene or propylene or a mixture thereof, m is 0 or 1, and n is from 1 to 20;

- ii) sulfoaroyl units of the formula -(O)C(C₆H₄)(SO₃-M⁺), wherein M is a salt forming cation;
- iii) modified poly(oxyethylene)oxy monoalkyl ether units of the formula R¹²O(CH₂CH₂O)_k-, wherein R¹² contains from 1 to 4 carbon atoms and k is from about 3 to about 100; and
- ethoxylated or propoxylated phenolsulfonate end-capping units of the formula MO₃S(C₆H₄)(OR¹³)_nO-, wherein n is from 1 to 20;
 M is a salt-forming cation; and R¹³ is ethylene, propylene and mixtures thereof.

This type of preferred non-cotton soil release polymer of the present invention may be described as having the formula

$$[(Cap)(R^4)_t][(A-R^1-A-R^2)_u(A-R^1-A-R^3)_v(A-R^1-A-R^5)_w$$

-A-R¹-A-][(R⁴)_t(Cap)]

wherein A is a carboxy linking moiety having the formula

R1 is arylene, preferably a 1,4-phenylene moiety having the formula

such that when A units and R^1 units are taken together in the formula A- R^1 -A they form a terephthalate unit having the formula

 R^2 units are ethyleneoxy or 1,2-propyleneoxy. R^2 units are combined with terephthalate moieties to form (A-R¹-A-R²) units having the formula

wherein R' and R" are either hydrogen or methyl provided that R' and R" are not both methyl at the same time.

R³ units are trifunctional, ester-forming, branching moieties having the formula

Preferably R^3 units comprise a glycerol moiety which is placed into the soil release polymer backbone to provide a branch point. When R^3 units are combined with terephthalate moieties to form units of the polymer backbone, for example, (A-R\frac{1}{2}-A-R\frac{3}{2}-A-R\frac{1}{2}-A units, these units have the formula

or the formula

wherein one terephthalate residue is taken to be a part of the $(A-R^1-A-R^3)$ unit while the second terephthalate comprises a part of another backbone unit, such as a $(A-R^1-A-R^2)$ unit, a $(A-R^1-A-R^3)$ unit, a $-A-R^1-A-[(R^4)_t(Cap)]$ unit or a second $(A-R^1-A-R^3)$ unit. The third functional group, which is the beginning of the branching chain, is also typically bonded to a terephthalate residue also a part of a $(A-R^1-A-R^2)$ unit, a $(A-R^1-A-R^3)$ unit.

An example of a section of a soil release polymer containing a "trifunctional, ester-forming, branching moiety" R³ unit which comprises a glycerol unit, has the formula

R⁴ units are R², R³ or R⁵ units. R⁵ units are units having the formula

wherein R⁹ is C₂-C₆ linear alkylene, C₃-C₆ branched alkylene, and mixtures thereof; preferably R¹⁰ is independently selected from hydrogen or -L-SO₃-M⁺; wherein L is a side chain moiety selected from the group consisting of alkylene, oxyalkylene, alkyleneoxyalkylene, arylene, oxyarylene, alkyleneoxyarylene, poly(oxyalkylene), oxyalkyleneoxyarylene, poly(oxyalkylene)oxyarlyene, alkylenepoly(oxyalkylene), and mixtures thereof; M is hydrogen or a salt forming cation; i has the value of 0 or 1;

Each carbon atom of the R⁹ units is substituted by R¹⁰ units that are independently selected from hydrogen or -L-SO₃-M⁺, provided no more than one -L-SO₃-M⁺ units is attached to an R⁹ unit; L is a side chain connecting moiety selected from the group consisting of alkylene, oxyalkylene, alkyleneoxyalkylene, arylene, oxyarylene, alkyleneoxyarylene, poly(oxyalkylene), oxyalkyleneoxyarylene, poly(oxyalkylene), oxyalkylene), and mixtures thereof.

M is a cationic moiety selected from the group consisting of lithium, sodium, potassium, calcium, and magnesium, preferably sodium and potassium.

Preferred R^5 moieties are essentially R^{10} substituted C_2 - C_6 alkylene chains. The R^5 units comprise either one C_2 - C_6 alkylene chain substituted by one or more independently selected R^{10} moieties (preferred) or two C_2 - C_6 alkylene chains said alkylene chains joined by an ether oxygen linkage, each alkylene chain substituted by one or more independently selected R^{10} moieties, that is R^5 may comprise two separate R^9 units, each of which is substituted by one or more independently selected R^{10} moieties. Preferably only one carbon atom of each R^9 moiety is substituted by an -L- SO_3 -M+ unit with the remaining R^{10} substituents comprising a hydrogen atom. When

the value of the index i is equal to 1 (two R^9 units comprise the R^5 unit), a preferred formula is

wherein each R⁹ comprises a C₂ alkylene moiety. Preferably one R¹⁰ moiety is -L-SO₃-M⁺, preferably the C₂ carbon is substituted by the -L-SO₃-M⁺ moiety, and the balance are hydrogen atoms, having therefore a formula:

wherein L is a polyethyleneoxymethyl substituent, x is from 0 to about 20.

As used herein, the term "R⁵ moieties consist essentially of units

having the index i equal to 0 wherein R^{10} units are hydrogen and one R^{10} units is equal to -L-SO₃-M⁺, wherein L is a side chain connecting moiety selected from the group consisting of alkylene, alkenylene, alkoxyalkylene, oxyalkylene, arylene, alkylarylene, alkoxyarylene and mixtures thereof", refers to the preferred compounds of the present invention wherein the R^{10} moieties consist of one -L-SO₃-M⁺ moiety and the rest of the R^{10} moieties are hydrogen atoms, for example a

which is capable of inclusion into the polymeric backbone of the soil release polymers of the present invention as an -A-R⁵-A- backbone segment. The units are easily incorporated into the oligomer or polymer backbone by using starting materials having the general formula

wherein x, for the purposes of the L moiety of the present invention, is from 0 to 20.

Other suitable monomers capable of inclusion into the backbone of the type A preferred non-cotton soil release polymers of the present invention as R⁵ moieties includes the alkylene poly(oxyalkylene)oxyarylene containing monomer having the general formula

wherein x is 0 to 20. A further example of a preferred monomer resulting in a preferred R⁵ unit wherein i is equal to 0, are the sodiosulfopoly(ethyleneoxy)methyl-1,2-propanediols having the formula

wherein x is from 0 to about 20; more preferred are the monomers

The preferred non-cotton soil release agents of the present invention in addition to the afore-mentioned R^1 , R^2 , R^3 , R^4 , and R^5 units also comprise one or more capping groups, -(Cap). The capping groups are independently selected from ethoxylated or propoxylated hydroxyethane and propanesulfonate units of the formula $(MO_3S)(CH_2)_m(R^{11}O)_n$ -, where M is a salt forming cation such as sodium or tetralkylammonium as described herein above, R^{11} is ethylene or propylene or a mixture thereof, m is 0 or 1, and n is from 1 to 20, preferably n is from 1 to about 4; sulfoaroyl units of the formula - $(O)C(C_6H_4)(SO_3^-M^+)$, wherein M is a salt forming cation as described herein above; modified poly(oxyethylene)oxy monoalkyl ether units of the formula $R^{12}O(CH_2CH_2O)_k$ - wherein R^{12} contains from 1 to 4 carbon atoms, R^{12} is preferably methyl, and k is from about 3 to about 100, preferably about 3 to about 50, more preferably 3 to about 30; and ethoxylated or propoxylated phenolsulfonate end-capping units of the formula $MO_3S(C_6H_4)(OR^{13})_nO$ -, wherein n is from to 20; M is a salt-forming cation; and R^{13} is ethylene, propylene and mixtures thereof.

Most preferred end capping unit is the isethionate-type end capping unit which is a hydroxyethane moiety, $(MO_3S)(CH_2)_m(R^{11}O)_n$, preferably R^{11} is ethyl, m is equal to 0, and n is from 2 to 4.

The value of t is 0 or 1; the value of u is from about 0 to about 60; the value of v is from about 0 to about 35; the value of w is from 0 to 35.

Preferred non-cotton soil release polymers of the present invention having the formula

$$[(Cap)(R^4)_t][(A-R^1-A-R^2)_u(A-R^1-A-R^3)_v(A-R^1-A-R^5)_w$$

-A-R¹-A-][(R⁴)_t(Cap)]

can be conveniently expressed as the following generic structural formula

$$\left[\begin{array}{c} O \\ O \\ C \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ C \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ O \\ C \end{array} \right] = \left[\begin{array}{c} O \\ O \\ C \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ C \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ C \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ C \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ O \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ O \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ O \\ O \end{array} \right] = \left[\begin{array}{c} O \\ O \\ O \end{array} \right] = \left[$$

The following structure is an example of the preferred non-cotton soil release polymers of the present invention.

$$NaO_{2}S(CH_{2}CH_{2}O)_{2-3}CH_{2}CH_{2} - OCH_{2}C$$

The above-described preferred non-cotton soil release agents are fully described in U.S. Patent Application Serial No. 08/545,351 filed November 22, 1995 which is a continuation-in-part of U.S. Patent Application Serial No. 08/355,938 filed December 14, 1994, both of which are incorporated herein by reference. Other non-cotton soil release polymers suitable for use in the compositions of the present invention are further described herein below.

The preferred non-cotton SRA's can be further described as oligomeric esters comprising: (1) a backbone comprising (a) at least one unit selected from the group consisting of dihydroxysulfonates, polyhydroxy sulfonates, a unit which is at least trifunctional whereby ester linkages are formed resulting in a branched oligomer backbone, and combinations thereof; (b) at least one unit which is a terephthaloyl moiety; and (c) at least one unsulfonated unit which is a 1,2-oxyalkyleneoxy moiety; and (2) one or more capping units selected from nonionic capping units, anionic capping units such as alkoxylated, preferably ethoxylated, isethionates, alkoxylated propanesulfonates, alkoxylated propanedisulfonates, alkoxylated phenolsulfonates, sulfoaroyl derivatives and mixtures thereof. Preferred are esters of the empirical formula:

{(CAP)x(EG/PG)y'(DEG)y"(PEG)y"'(T)z(SIP)z'(SEG)q(B)m} wherein CAP, EG/PG, PEG, T and SIP are as defined as terephthaloyl (T), sulfoisophthaloyl (SIP), oxyethyleneoxy and oxy-1,2-propylene (EG/PG) units, end-caps (CAP), poly(ethyleneglycol) (PEG), (DEG) represents di(oxyethylene)oxy units, (SEG) represents units derived from the sulfoethyl ether of glycerin and related moiety units, (B) represents branching units which are at least trifunctional whereby ester linkages are formed resulting in a branched oligomer backbone, x is from about 1 to about 12, y' is from about 0.5 to about 25, y" is from 0 to about 12, y" is from 0 to about 10, y'+y"+y" totals from about 0.5 to about 25, z is from about 1.5 to about 25, z' is from 0 to about 12; z + z' totals from about 1.5 to about 25, q is from about 0.05 to about 12; m is from about 0.01 to about 10, and x, y', y", y"', z, z', q and m represent the average number of moles of the corresponding units per mole of said ester and said ester has a molecular weight ranging from about 500 to about 5,000.

Preferred SEG and CAP monomers for the above esters include Na-2-(2-,3-dihydroxypropoxy)ethanesulfonate ("SEG"), Na-2-{2-(2-hydroxyethoxy) ethoxy} ethanesulfonate ("SE3") and its homologs and mixtures thereof and the products of ethoxylating and sulfonating allyl alcohol. Preferred SRA esters in this class include the product of transesterifying and oligomerizing sodium 2-{2-(2-hydroxyethoxyet

Preferred non-cotton soil release agent - B. A second preferred class of suitable SRA's include a sulfonated product of a substantially linear ester oligomer comprised of an oligomeric ester backbone of terephthaloyl and oxyalkyleneoxy repeat units and allylderived sulfonated terminal moieties covalently attached to the backbone Such ester oligomers can be prepared by: (a) ethoxylating allyl alcohol; (b) reacting the product of (a) with dimethyl terephthalate ("DMT") and 1,2-propylene glycol ("PG") in a two-stage transesterification/oligomerization procedure; and (c) reacting the product of (b) with sodium metabisulfite in water.

Suitable for use in the laundry detergent compositions of the present invention are preferred non-cotton soil release polymers comprising:

- a) one or two terminal units selected from the group consisting of
 - i) -(CH₂)_q(CHSO₃M)CH₂SO₃M,
 - ii) -(CH₂)_q(CHSO₂M)CH₂SO₃M,
 - iii) -CH2CH2SO3M,
 - iv) and mixtures thereof; wherein q has the value from 1 to about 4, M is a water soluble cation, preferably sodium;
- b) a backbone comprising:
 - i) arylene units, preferably terephthalate units having the formula:

ii) ethyleneoxy units having the formula:

wherein the value of n is from about 1 to about 20; and

iii) 1,2-propyleneoxy units having the formula:

wherein the value of n is from about 1 to about 20, and wherein further the preferred backbone of this preferred non-cotton soil release polymer has a backbone comprising arylene repeat units which alternate with the ethyleneoxy and 1,2-propyleneoxy units, such that the mole ratio of ethyleneoxy to 1,2-propyleneoxy units is from 0:1 to about 0.9:0.1, preferably from about 0:1 to about 0.4:0.6, more preferably the arylene units alternate with essentially 1,2-propyleneoxy units.

However, other combinations of the above-identified units may be used to form non-cotton soil release polymers suitable for use in the compositions of the present invention. These combinations are more thoroughly described in U.S. Patent 4,968,451, Scheibel et al., issued November 6, 1990 and incorporated herein by reference.

<u>Preferred non-cotton soil release agent - C</u>. Suitable for use in the laundry detergent compositions of the present invention are preferred non-cotton soil release polymers having the formula

$$(Cap)[(A-R^1-A-R^2)_{11}(A-R^3-A-R^2)_{12}-A-R^4-A-](Cap)$$

wherein A is a carboxy linking moiety, preferably A is a carboxy linking moiety having the formula

R¹ is an arylene moiety, preferably 1,4-phenylene moiety having the formula

wherein for R¹ moieties, the degree of partial substitution with arylene moieties other than 1,4-phenylene should be such that the soil release properties of the compound are not adversely affected to any great extent. Generally, the partial substitution which can be tolerated will depend upon the backbone length of the compound.

 R^2 moieties are ethylene moieties or substituted ethylene moieties having C_1 - C_4 alkyl or alkoxy substituents. As used herein, the term "the R^2 moieties are essentially ethylene moieties or substituted ethylene moieties having C_1 - C_4 alkyl or alkoxy substituents" refers to compounds of the present invention where the R^2 moieties consist entirely of ethylene or substituted ethylene moieties or a partially substituted with other compatible moieties. Examples of these other moieties include 1,3-propylene, 1,4-butylene, 1,5-pentylene, or 1,6-hexylene, 1,2-hydroxyalkylenes and oxyalkylenes.

For the R² moieties, the degree of partial substitution with these other moieties should be such that the soil release properties of the compounds are not adversely affected to any great extent. For example, for polyesters made according to the present invention with a 75:25 mole ratio of diethylene glycol (-CH₂CH₂OCH₂CH₂-) to ethylene glycol (ethylene) have adequate soil release activity.

For the R³ moieties, suitable substituted C₂-C₁₈ hydrocarbylene moieties can include substituted C₂-C₁₂ alkylene, alkenylene, arylene, alkarylene and like moieties,

The substituted alkylene or alkenylene moieties can be linear, branched or cyclic. Also, the R^3 can all be the same (e.g. all substituted arylene) or a mixture (e.g. a mixture of substituted arylenes and substituted alkylenes). Preferred R^3 moieties are those which are substituted 1,3-phenylene, preferably 5-sulfo-1,3-phenylene. R^3 moieties are also - $A-[(R^2-A-R^4)]$ -Cap wherein R^4 is R^1 , R^3 , and mixtures thereof.

The preferred (Cap) moieties comprise units having the formula

$$--[(R5O)m(CH2CH2O)n]X$$

wherein R^5 is C_1 - C_4 alkylene, or the moiety - R^2 -A- R^6 - wherein R^6 is C_2 - C_{12} alkylene, alkenylene, arylene or alkarylene moiety, X is C_1 - C_4 alkyl, preferably methyl; the indices m and n are such that the moiety - CH_2CH_2O - comprises at least 50% by weight of the moiety

$$-[(R^5O)_m(CH_2CH_2O)_n]X$$

provided that when R^5 is the moiety $-R^2$ -A-R⁶-, m is at least 1; each n is at least about 10, the indices u and v are such that the sum of u + v is from about 3 to about 25; the index w is 0 or at least 1; and when w is at least 1, the indices u, v and w have the values such that the sum of u + v + w is from about 3 to about 25.

An example of this type of non-cotton soil release block polyester has the formula

wherein the R^2 moieties are essentially ethylene moieties, 1,2-propylene moieties, and mixtures thereof; the R^3 moieties are all potassium or preferably sodium 5-sulfo-1,3-phenylene moieties; the R^4 moieties are R^1 or R^3 moieties, or mixtures thereof; each X is ethyl, methyl, preferably methyl; each n is from about 12 to about 43; when w is 0, u + v is from about 3 to about 10; when w is at least 1, u + v + w is from about 3 to about 10.

The above non-cotton soil release polymers of the formula

$$(Cap)[(A-R^1-A-R^2)_u(A-R^3-A-R^2)_v-A-R^4-A-](Cap)$$

are further described in detail in U.S. Patent 4,702,857, Gosselink, issued October 27, 1987 and incorporated herein by reference.

In addition to the above-described non-cotton soil release polymers, other soil release polymers suitable for use in the liquid laundry detergent compositions of the present invention are further described herein below.

Any other anionic non-cotton soil release agent is suitable for use in the compositions of the present invention alone or in combination except for carboxymethylcellulose (CMC) which cannot be used alone. If the formulator selects CMC for use as an anionic soil release agent in the laundry detergent compositions of the present invention, carboxymethylcellulose must be present in an amount greater than 0.2% by weight, of the composition.

Cotton Soil Release Polymers

The cotton soil release agents useful in the present invention liquid detergent compositions are water-soluble or dispersible, modified polyamines. These polyamines comprise backbones that can be either linear or cyclic. The polyamine backbones can also comprise polyamine branching chains to a greater or lesser degree. In general, the polyamine backbones described herein are modified in such a manner that each nitrogen of the polyamine chain is thereafter described in terms of a unit that is substituted, quaternized, oxidized, or combinations thereof.

For the purposes of the present invention the term "modification" is defined as replacing a backbone -NH hydrogen atom by an E unit (substitution), quaternizing a backbone nitrogen (quaternized) or oxidizing a backbone nitrogen to the N-oxide (oxidized). The terms "modification" and "substitution" are used interchangably when referring to the process of replacing a hydrogen atom attached to a backbone nitrogen with an E unit. Quaternization or oxidation may take place in some circumstances without substitution, but substitution is preferably accompanied by oxidation or quaternization of at least one backbone nitrogen.

The linear or non-cyclic polyamine backbones that comprise the cotton soil release agents of the present invention have the general formula:

$$H_{2N-R}$$
 $-R_{1n-1}$ $-[N-R]_{m}$ $-[N-R]_{n}$ $-NH_{2}$

said backbones prior to subsequent modification, comprise primary, secondary and tertiary amine nitrogens connected by R "linking" units. The cyclic polyamine backbones comprising the cotton soil release agents of the present invention have the general formula:

$$[H_2N-R]_{m'k+1}$$
 $[N-R]_m$ $[N-R]_k-NH_2$

said backbones prior to subsequent modification, comprise primary, secondary and tertiary amine nitrogens connected by R "linking" units

For the purpose of the present invention, primary amine nitrogens comprising the backbone or branching chain once modified are defined as V or Z "terminal" units. For example, when a primary amine moiety, located at the end of the main polyamine backbone or branching chain having the structure

is modified according to the present invention, it is thereafter defined as a V "terminal" unit, or simply a V unit. However, for the purposes of the present invention, some or all of the primary amine moieties can remain unmodified subject to the restrictions further described herein below. These unmodified primary amine moieties by virtue of their position in the backbone chain remain "terminal" units. Likewise, when a primary amine moiety, located at the end of the main polyamine backbone having the structure

is modified according to the present invention, it is thereafter defined as a Z "terminal" unit, or simply a Z unit. This unit can remain unmodified subject to the restrictions further described herein below.

In a similar manner, secondary amine nitrogens comprising the backbone or branching chain once modified are defined as W "backbone" units. For example, when a secondary amine moiety, the major constituent of the backbones and branching chains of the present invention, having the structure

is modified according to the present invention, it is thereafter defined as a W "backbone" unit, or simply a W unit. However, for the purposes of the present invention, some or all of the secondary amine moieties can remain unmodified. These unmodified secondary amine moieties by virtue of their position in the backbone chain remain "backbone" units.

In a further similar manner, tertiary amine nitrogens comprising the backbone or branching chain once modified are further referred to as Y "branching" units. For example, when a tertiary amine moiety, which is a chain branch point of either the polyamine backbone or other branching chains or rings, having the structure

is modified according to the present invention, it is thereafter defined as a Y "branching" unit, or simply a Y unit. However, for the purposes of the present invention, some or all

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or the tertiary amine moieties can remain unmodified. These unmodified tertiary amine moieties by virtue of their position in the backbone chain remain "branching" units. The R units associated with the V, W and Y unit nitrogens which serve to connect the polyamine nitrogens, are described herein below.

The final modified structure of the polyamines of the present invention can be therefore represented by the general formula

$$V_{(n+1)}W_mY_nZ$$

for linear polyamine cotton soil release polymers and by the general formula

$$V_{(n-k+1)}W_mY_nY_kZ$$

for cyclic polyamine cotton soil release polymers. For the case of polyamines comprising rings, a Y' unit of the formula

serves as a branch point for a backbone or branch ring. For every Y' unit there is a Y unit having the formula

that will form the connection point of the ring to the main polymer chain or branch. In the unique case where the backbone is a complete ring, the polyamine backbone has the formula

$$[H_2N-R]_n$$
- $[N-R]_m$ - $[N-R]_n$ -

therefore comprising no Z terminal unit and having the formula

$$V_{n-k}W_mY_nY_k'$$

wherein k is the number of ring forming branching units. Preferably the polyamine backbones of the present invention comprise no rings.

In the case of non-cyclic polyamines, the ratio of the index n to the index m relates to the relative degree of branching. A fully non-branched linear modified polyamine according to the present invention has the formula

$$VW_mZ$$

that is, n is equal to 0. The greater the value of n (the lower the ratio of m to n), the greater the degree of branching in the molecule. Typically the value for m ranges from a minimum value of 4 to about 400, however larger values of m, especially when the value of the index n is very low or nearly 0, are also preferred.

Each polyamine nitrogen whether primary, secondary or tertiary, once modified according to the present invention, is further defined as being a member of one of three general classes; simple substituted, quaternized or oxidized. Those polyamine nitrogen units not modified are classed into V, W, Y, or Z units depending on whether they are primary, secondary or tertiary nitrogens. That is unmodified primary amine nitrogens are V or Z units, unmodified secondary amine nitrogens are W units and unmodified tertiary amine nitrogens are Y units for the purposes of the present invention.

Modified primary amine moieties are defined as V "terminal" units having one of three forms:

a) simple substituted units having the structure:

b) quaternized units having the structure:

wherein X is a suitable counter ion providing charge balance; and

c) oxidized units having the structure:

Modified secondary amine moieties are defined as W "backbone" units having one of three forms:

a) simple substituted units having the structure:

b) quaternized units having the structure:

wherein X is a suitable counter ion providing charge balance; and

c) oxidized units having the structure:

Modified tertiary amine moieties are defined as Y "branching" units having one of three forms:

a) unmodified units having the structure:

b) quaternized units having the structure:

wherein X is a suitable counter ion providing charge balance; and

c) oxidized units having the structure:

Certain modified primary amine moieties are defined as Z "terminal" units having one of three forms:

a) simple substituted units having the structure:

b) quaternized units having the structure:

wherein X is a suitable counter ion providing charge balance; and

c) oxidized units having the structure:



When any position on a nitrogen is unsubstituted of unmodified, it is understood that hydrogen will substitute for E. For example, a primary amine unit comprising one E unit in the form of a hydroxyethyl moiety is a V terminal unit having the formula (HOCH₂CH₂)HN-.

For the purposes of the present invention there are two types of chain terminating units, the V and Z units. The Z "terminal" unit derives from a terminal primary amino moiety of the structure -NH₂. Non-cyclic polyamine backbones according to the present invention comprise only one Z unit whereas cyclic polyamines can comprise no Z units. The Z "terminal" unit can be substituted with any of the E units described further herein below, except when the Z unit is modified to form an N-oxide. In the case where the Z unit nitrogen is oxidized to an N-oxide, the nitrogen must be modified and therefore E cannot be a hydrogen.

The polyamines of the present invention comprise backbone R "linking" units that serve to connect the nitrogen atoms of the backbone. R units comprise units that for the purposes of the present invention are referred to as "hydrocarbyl R" units and "oxy R" units. The "hydrocarbyl" R units are C₂-C₁₂ alkylene, C₄-C₁₂ alkenylene, C₃-C₁₂ hydroxyalkylene wherein the hydroxyl moiety may take any position on the R unit chain except the carbon atoms directly connected to the polyamine backbone nitrogens; C₄-C₁₂ dihydroxyalkylene wherein the hydroxyl moieties may occupy any two of the carbon atoms of the R unit chain except those carbon atoms directly connected to the polyamine backbone nitrogens; C₈-C₁₂ dialkylarylene which for the purpose of the present invention are arylene moieties having two alkyl substituent groups as part of the linking chain. For example, a dialkylarylene unit has the formula

$$-(CH_2)_2$$
 $-(CH_2)_4$ $-(CH_2)_4$

although the unit need not be 1,4-substituted, but can also be 1,2 or 1,3 substituted C_2 - C_{12} alkylene, preferably ethylene, 1,2-propylene, and mixtures thereof, more preferably ethylene. The "oxy" R units comprise - $(R^1O)_xR^5(OR^1)_{X^-}$, - $CH_2CH(OR^2)CH_2O)_z(R^1O)_yR^1(OCH_2CH(OR^2)CH_2)_{W^-}$, - $CH_2CH(OR^2)CH_2O)_z(R^1O)_yR^1(OCH_2CH(OR^2)CH_2)_{W^-}$, - $CH_2CH(OR^2)CH_2O)_z(R^1O)_z(R$

 R^1 units are C_2 - C_6 alkylene, and mixtures thereof, preferably ethylene. R^2 is hydrogen, and $-(R^1O)_xB$, preferably hydrogen.

 R^3 is C_1 - C_{18} alkyl, C_7 - C_{12} arylalkylene, C_7 - C_{12} alkyl substituted aryl, C_6 - C_{12} aryl, and mixtures thereof, preferably C_1 - C_{12} alkyl, C_7 - C_{12} arylalkylene, more preferably C_1 - C_{12} alkyl, most preferably methyl. R^3 units serve as part of E units described herein below.

R⁴ is C₁-C₁₂ alkylene, C₄-C₁₂ alkenylene, C₈-C₁₂ arylalkylene, C₆-C₁₀ arylene, preferably C₁-C₁₀ alkylene, C₈-C₁₂ arylalkylene, more preferably C₂-C₈ alkylene, most preferably ethylene or butylene.

 R^5 is C1-C12 alkylene, C3-C12 hydroxyalkylene, C4-C12 dihydroxyalkylene, C8-C12 dialkylarylene, -C(O)-, -C(O)NHR⁶NHC(O)-, -C(O)(R⁴)_rC(O)-, -R¹(OR¹)-, -CH2CH(OH)CH2O(R¹O)_yR¹OCH2CH(OH)CH2-, -C(O)(R⁴)_rC(O)-, -CH2CH(OH)CH2-, R⁵ is preferably ethylene, -C(O)-, -C(O)NHR⁶NHC(O)-, -R¹(OR¹)-, -CH2CH(OH)CH2-, -CH2CH(OH)CH2O(R¹O)_yR¹OCH2CH-(OH)CH2-, more preferably -CH2CH(OH)CH2-.

R⁶ is C₂-C₁₂ alkylene or C₆-C₁₂ arylene.

The preferred "oxy" R units are further defined in terms of the R¹, R², and R⁵ units. Preferred "oxy" R units comprise the preferred R¹, R², and R⁵ units. The preferred cotton soil release agents of the present invention comprise at least 50% R¹

units that are ethylene. Preferred R¹, R², and R⁵ units are combined with the "oxy" R units to yield the preferred "oxy" R units in the following manner.

- i) Substituting more preferred R⁵ into -(CH₂CH₂O)_xR⁵(OCH₂CH₂O)_x-yields -(CH₂CH₂O)_xCH₂CHOHCH₂(OCH₂CH₂O)_x-.
- ii) Substituting preferred R^1 and R^2 into -(CH₂CH(OR²)CH₂O)_z-(R¹O)_yR¹O(CH₂CH(OR²)CH₂)_w- yields -(CH₂CH(OH)CH₂O)_z-(CH₂CH₂O)_yCH₂CH₂O(CH₂CH(OH)CH₂)_w-.
- iii) Substituting preferred R² into -CH₂CH(OR²)CH₂- yields -CH₂CH(OH)CH₂-.

E units are selected from the group consisting of hydrogen, C_1 - C_{22} alkyl, C_3 - C_{22} alkenyl, C_7 - C_{22} arylalkyl, C_2 - C_{22} hydroxyalkyl, -(CH₂)_pCO₂M, -(CH₂)_qSO₃M, -CH(CH₂CO₂M)CO₂M, -(CH₂)_pPO₃M, -(R¹O)_mB, -C(O)R³, preferably hydrogen, C_2 - C_{22} hydroxyalkylene, benzyl, C_1 - C_{22} alkylene, -(R¹O)_mB, -C(O)R³, -(CH₂)_pCO₂M, -(CH₂)_qSO₃M, -CH(CH₂CO₂M)CO₂M, more preferably C_1 - C_{22} alkylene, -(R¹O)_xB, -C(O)R³, -(CH₂)_pCO₂M, -(CH₂)_qSO₃M, -CH(CH₂CO₂M)CO₂M, most preferably C_1 - C_{22} alkylene, -(R¹O)_xB, and -C(O)R³. When no modification or substitution is made on a nitrogen then hydrogen atom will remain as the moiety representing E.

E units do not comprise hydrogen atom when the V, W or Z units are oxidized, that is the nitrogens are N-oxides. For example, the backbone chain or branching chains do not comprise units of the following structure:

Additionally, E units do not comprise carbonyl moieties directly bonded to a nitrogen atom when the V, W or Z units are oxidized, that is, the nitrogens are N-oxides. According to the present invention, the E unit -C(O)R³ moiety is not bonded to an N-oxide modified nitrogen, that is, there are no N-oxide amides having the structure

nor combinations thereof.

B is hydrogen, C_1 - C_6 alkyl, -(CH₂)_qSO₃M, -(CH₂)_pCO₂M, -(CH₂)_q-(CHSO₃M)CH₂SO₃M, -(CH₂)_q(CHSO₂M)CH₂SO₃M, -(CH₂)_pPO₃M, -PO₃M, preferably hydrogen, -(CH₂)_qSO₃M, -(CH₂)_q(CHSO₃M)CH₂SO₃M, -(CH₂)_q-(CHSO₂M)CH₂SO₃M, more preferably hydrogen or -(CH₂)_qSO₃M.

M is hydrogen or a water soluble cation in sufficient amount to satisfy charge balance. For example, a sodium cation equally satisfies -(CH₂)_pCO₂M, and -(CH₂)_qSO₃M, thereby resulting in -(CH₂)_pCO₂Na, and -(CH₂)_qSO₃Na moieties. More than one monovalent cation, (sodium, potassium, etc.) can be combined to satisfy the required chemical charge balance. However, more than one anionic group may be charge balanced by a divalent cation, or more than one mono-valent cation may be necessary to satisfy the charge requirements of a poly-anionic radical. For example, a -(CH₂)_pPO₃M moiety substituted with sodium atoms has the formula -(CH₂)_pPO₃Na₃. Divalent cations such as calcium (Ca²⁺) or magnesium (Mg²⁺) may be substituted for or combined with other suitable mono-valent water soluble cations. Preferred cations are sodium and potassium, more preferred is sodium.

X is a water soluble anion such as chlorine (Cl⁻), bromine (Br⁻) and iodine (l⁻) or X can be any negatively charged radical such as sulfate (SO₄²-) and methosulfate (CH₃SO₃⁻).

The formula indices have the following values: p has the value from 1 to 6, q has the value from 0 to 6; r has the value 0 or 1; w has the value 0 or 1, x has the value from 1 to 100; y has the value from 0 to 100; z has the value 0 or 1; m has the value from 4 to about 400, n has the value from 0 to about 200; m + n has the value of at least 5.

The preferred cotton soil release agents of the present invention comprise polyamine backbones wherein less than about 50% of the R groups comprise "oxy" R units, preferably less than about 20%, more preferably less than 5%, most preferably the R units comprise no "oxy" R units.

The most preferred cotton soil release agents which comprise no "oxy" R units comprise polyamine backbones wherein less than 50% of the R groups comprise more than 3 carbon atoms. For example, ethylene, 1,2-propylene, and 1,3-propylene comprise 3 or less carbon atoms and are the preferred "hydrocarby!" R units. That is when

backbone R units are C₂-C₁₂ alkylene, preferred is C₂-C₃ alkylene, most preferred is ethylene.

The cotton soil release agents of the present invention comprise modified homogeneous and non-homogeneous polyamine backbones, wherein 100% or less of the -NH units are modified. For the purpose of the present invention the term "homogeneous polyamine backbone" is defined as a polyamine backbone having R units that are the same (i.e., all ethylene). However, this sameness definition does not exclude polyamines that comprise other extraneous units comprising the polymer backbone which are present due to an artifact of the chosen method of chemical synthesis. For example, it is known to those skilled in the art that ethanolamine may be used as an "initiator" in the synthesis of polyethyleneimines, therefore a sample of polyethyleneimine that comprises one hydroxyethyl moiety resulting from the polymerization "initiator" would be considered to comprise a homogeneous polyamine backbone for the purposes of the present invention. A polyamine backbone comprising all ethylene R units wherein no branching Y units are present is a homogeneous backbone. A polyamine backbone comprising all ethylene R units is a homogeneous backbone regardless of the degree of branching or the number of cyclic branches present.

For the purposes of the present invention the term "non-homogeneous polymer backbone" refers to polyamine backbones that are a composite of various R unit lengths and R unit types. For example, a non-homogeneous backbone comprises R units that are a mixture of ethylene and 1,2-propylene units. For the purposes of the present invention a mixture of "hydrocarbyl" and "oxy" R units is not necessary to provide a non-homogeneous backbone. The proper manipulation of these "R unit chain lengths" provides the formulator with the ability to modify the solubility and fabric substantivity of the cotton soil release agents of the present invention.

Preferred cotton soil release polymers of the present invention comprise homogeneous polyamine backbones that are totally or partially substituted by polyethyleneoxy moieties, totally or partially quaternized amines, nitrogens totally or partially oxidized to N-oxides, and mixtures thereof. However, not all backbone amine nitrogens must be modified in the same manner, the choice of modification being left to the specific needs of the formulator. The degree of ethoxylation is also determined by the specific requirements of the formulator.

The preferred polyamines that comprise the backbone of the compounds of the present invention are generally polyalkyleneamines (PAA's), polyalkyleneimines (PAI's), preferably polyethyleneamine (PEA's), polyethyleneimines (PEI's), or PEA's or PEI's connected by moieties having longer R units than the parent PAA's, PAI's, PEA's or

PEI's. A common polyalkyleneamine (PAA) is tetrabutylenepentamine. PEA's are obtained by reactions involving ammonia and ethylene dichloride, followed by fractional distillation. The common PEA's obtained are triethylenetetramine (TETA) and teraethylenepentamine (TEPA). Above the pentamines, i.e., the hexamines, heptamines, octamines and possibly nonamines, the cogenerically derived mixture does not appear to separate by distillation and can include other materials such as cyclic amines and particularly piperazines. There can also be present cyclic amines with side chains in which nitrogen atoms appear. See U.S. Patent 2,792,372, Dickinson, issued May 14, 1957, which describes the preparation of PEA's.

Preferred amine polymer backbones comprise R units that are C₂ alkylene (ethylene) units, also known as polyethylenimines (PEI's). Preferred PEI's have at least moderate branching, that is the ratio of m to n is less than 4:1, however PEI's having a ratio of m to n of about 2:1 are most preferred. Preferred backbones, prior to modification have the general formula:

wherein m and n are the same as defined herein above. Preferred PEI's, prior to modification, will have a molecular weight greater than about 200 daltons.

The relative proportions of primary, secondary and tertiary amine units in the polyamine backbone, especially in the case of PEI's, will vary, depending on the manner of preparation. Each hydrogen atom attached to each nitrogen atom of the polyamine backbone chain represents a potential site for subsequent substitution, quaternization or oxidation.

These polyamines can be prepared, for example, by polymerizing ethyleneimine in the presence of a catalyst such as carbon dioxide, sodium bisulfite, sulfuric acid, hydrogen peroxide, hydrochloric acid, acetic acid, etc. Specific methods for preparing these polyamine backbones are disclosed in U.S. Patent 2,182,306, Ulrich et al., issued December 5, 1939; U.S. Patent 3,033,746, Mayle et al., issued May 8, 1962; U.S. Patent 2,208,095, Esselmann et al., issued July 16, 1940; U.S. Patent 2,806,839, Crowther, issued September 17, 1957; and U.S. Patent 2,553,696, Wilson, issued May 21, 1951; all herein incorporated by reference.

Examples of modified cotton soil release polymers of the present invention comprising PEI's, are illustrated in Formulas I - V:

Formula I depicts a preferred cotton soil release polymer comprising a PEI backbone wherein all substitutable nitrogens are modified by replacement of hydrogen with a polyoxyalkyleneoxy unit, -(CH₂CH₂O)₂₀H, having the formula:

$$[H(OCH_{2}CH_{2})_{20}]_{2}N \longrightarrow N[(CH_{2}CH_{2}O)_{20}H]_{2}$$

$$H(OCH_{2}CH_{2})_{20} \longrightarrow (CH_{2}CH_{2}O)_{20}H$$

$$(CH_{2}CH_{2}O)_{20}H \longrightarrow N[(CH_{2}CH_{2}O)_{20}H]_{2}$$

$$(CH_{2}CH_{2}O)_{20}H \longrightarrow N[(CH_{2}CH_{2}O)_{20}H]_{2}$$

$$(CH_{2}CH_{2}O)_{20}H$$

$$(CH_{2}CH_{2}O)_{20}H]_{2}$$

Formula I

Formula II depicts a cotton soil release polymer comprising a PEI backbone wherein all substitutable nitrogens are modified by replacement of hydrogen with a polyoxyalkyleneoxy unit, -(CH₂CH₂O)₇H, having the formula

$$[H(OCH_{2}CH_{2})_{7}]_{2}N \\ \qquad \qquad \qquad N[(CH_{2}CH_{2}O)_{7}H]_{2} \\ \qquad \qquad \qquad \\ (CH_{2}CH_{2}O)_{7}H \\ \qquad \qquad \\ (CH_{2}CH_{2}O)_{7}$$

Formula II

This is an example of a cotton soil release polymer that is fully modified by one type of moiety.

Formula III depicts a cotton soil release polymer comprising a PEI backbone wherein all substitutable primary amine nitrogens are modified by replacement of hydrogen with a polyoxyalkyleneoxy unit, -(CH₂CH₂O)₇H, the molecule is then modified by subsequent oxidation of all oxidizable primary and secondary nitrogens to N-oxides, said cotton soil release agent having the formula

Formula III

Formula IV depicts a cotton soil release polymer comprising a PEI backbone wherein all backbone hydrogen atoms are substituted and some backbone amine units are quaternized. The substituents are polyoxyalkyleneoxy units, -(CH₂CH₂O)₇H, or methyl groups. The modified PEI cotton soil release polymer has the formula

Formula IV

Formula V depicts a cotton soil release polymer comprising a PEI backbone wherein the backbone nitrogens are modified by substitution (i.e. by -(CH₂CH₂O)₇H or methyl), quaternized, oxidized to N-oxides or combinations thereof. The resulting cotton soil release polymer has the formula

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Formula V

In the above examples, not all nitrogens of a unit class comprise the same modification. The present invention allows the formulator to have a portion of the secondary amine nitrogens ethoxylated while having other secondary amine nitrogens oxidized to N-oxides. This also applies to the primary amine nitrogens, in that the formulator may choose to modify all or a portion of the primary amine nitrogens with one or more substituents prior to oxidation or quaternization. Any possible combination of E groups can be substituted on the primary and secondary amine nitrogens, except for the restrictions described herein above.

The laundry detergent compositions according to the present invention comprise adjunct ingredients and carriers, said adjunct ingredients are selected from the group consisting of builders, optical brighteners, bleaches, bleach boosters, bleach activators, other non-cotton soil release polymers, dye transfer agents, dispersents, enzymes, enzyme activators, suds suppressors, dyes, perfumes, colorants, filler salts, hydrotropes, and mixtures thereof, however this list is not meant to be exhaustive or to exclude any suitable material used by the formulator.

Detersive surfactants

In addition to the anionic and nonionic detersive surfactants described herein above, other detersive surfactants that are suitable for use in the present invention are cationic, anionic, nonionic, ampholytic, zwitterionic, and mixtures thereof, further described herein below.

Nonlimiting examples of other surfactants useful herein typically at levels from about 1% to about 55%, by weight, include the conventional C11-C18 alkyl benzene sulfonates ("LAS"), the C10-C18 secondary (2,3) alkyl sulfates of the formula CH₃(CH₂)_x(CHOSO₃⁻M⁺) CH₃ and CH₃ (CH₂)_y(CHOSO₃⁻M⁺) CH₂CH₃ where x and (y + 1) are integers of at least about 7, preferably at least about 9, and M is a water-solubilizing cation, especially sodium, unsaturated sulfates such as oleyl sulfate,

C₁₀-C₁₈ alkyl alkoxy carboxylates (especially the EO 1-5 ethoxycarboxylates), the C₁₀₋₁₈ glycerol ethers, the C₁₀-C₁₈ alkyl polyglycosides and their corresponding sulfated polyglycosides, and C₁₂-C₁₈ alpha-sulfonated fatty acid esters. If desired, the conventional nonionic and amphoteric surfactants such as the C₁₂-C₁₈ alkyl ethoxylates ("AE") including the so-called narrow peaked alkyl ethoxylates and C₆-C₁₂ alkyl phenol alkoxylates (especially ethoxylates and mixed ethoxy/propoxy), C₁₂-C₁₈ betaines and sulfobetaines ("sultaines"), C₁₀-C₁₈ amine oxides, and the like, can also be included in the overall compositions. The C₁₀-C₁₈ N-alkyl polyhydroxy fatty acid amides can also be used. Typical examples include the C₁₂-C₁₈ N-methylglucamides. See WO 9,206,154. Other sugar-derived surfactants include the N-alkoxy polyhydroxy fatty acid amides, such as C₁₀-C₁₈ N-(3-methoxypropyl) glucamide. C₁₀-C₂₀ conventional soaps may also be used. If high sudsing is desired, the branched-chain C₁₀-C₁₆ soaps may be used. Mixtures of anionic and nonionic surfactants are especially useful. Other conventional useful surfactants are listed in standard texts.

Other anionic surfactants useful for detersive purposes can also be included in the compositions hereof. These can include salts (including, for example, sodium potassium, ammonium, and substituted ammonium salts such a mono-, di- and triethanolamine salts) of soap, C9-C20 linear alkylbenzenesulphonates, C8-C22 primary or secondary alkanesulphonates, C8-C24 olefinsulphonates, sulphonated polycarboxylic acids, alkyl glycerol sulfonates, fatty acyl glycerol sulfonates, fatty oleyl glycerol sulfates, alkyl phenol ethylene oxide ether sulfates, paraffin sulfonates, alkyl phosphates, isothionates such as the acyl isothionates, N-acyl taurates, fatty acid amides of methyl tauride, alkyl succinamates and sulfosuccinates, monoesters of sulfosuccinate (especially saturated and unsaturated C₁₂-C₁₈ monoesters) diesters of sulfosuccinate (especially saturated and unsaturated C6-C14 diesters), N-acyl sarcosinates, sulfates of alkylpolysaccharides such as the sulfates of alkylpolyglucoside, branched primary alkyl sulfates, alkyl polyethoxy carboxylates such as those of the formula RO(CH2CH2O)kCH2COO-M+ wherein R is a C8-C22 alkyl, k is an integer from 0 to 10, and M is a soluble salt-forming cation, and fatty acids esterified with isethionic acid and neutralized with sodium hydroxide. Further examples are given in Surface Active Agents and Detergents (Vol. I and II by Schwartz, Perry and Berch).

Non-cotton Soil Release Agent

Known polymeric soil release agents, hereinafter "SRA", can optionally be employed in the present detergent compositions. If utilized, SRA's will generally comprise from 0.01% to 10.0%, typically from 0.1% to 5%, preferably from 0.2% to 3.0% by weight, of the compositions.

Preferred SRA's typically have hydrophilic segments to hydrophilize the surface of hydrophobic fibers such as polyester and nylon, and hydrophobic segments to deposit upon hydrophobic fibers and remain adhered thereto through completion of washing and rinsing cycles, thereby serving as an anchor for the hydrophilic segments. This can enable stains occurring subsequent to treatment with the SRA to be more easily cleaned in later washing procedures.

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SRA's can include a variety of charged, e.g., anionic or even cationic species, see U.S. 4,956,447, issued September 11, 1990 to Gosselink, et al., as well as noncharged monomer units, and their structures may be linear, branched or even star-shaped. They may include capping moieties which are especially effective in controlling molecular weight or altering the physical or surface-active properties. Structures and charge distributions may be tailored for application to different fiber or textile types and for varied detergent or detergent additive products.

Preferred SRA's include oligomeric terephthalate esters, typically prepared by processes involving at least one transesterification/oligomerization, often with a metal catalyst such as a titanium(IV) alkoxide. Such esters may be made using additional monomers capable of being incorporated into the ester structure through one, two, three, four or more positions, without, of course, forming a densely crosslinked overall structure.

Other SRA's include the nonionic end-capped 1,2-propylene/polyoxyethylene terephthalate polyesters of U.S. 4,711,730, December 8, 1987 to Gosselink et al., for example those produced by transesterification/oligomerization of poly(ethyleneglycol) methyl ether, DMT, PG and poly(ethyleneglycol) ("PEG"). Other examples of SRA's include: the partly- and fully- anionic-end-capped oligomeric esters of U.S. 4,721,580, January 26, 1988 to Gosselink, such as oligomers from ethylene glycol ("EG"), PG, DMT and Na-3,6-dioxa-8-hydroxyoctanesulfonate; and the anionic, especially sulfoaroyl, end-capped terephthalate esters of U.S. 4,877,896, October 31, 1989 to Maldonado, the latter being typical of SRA's useful in both laundry and fabric conditioning products, an example being an ester composition made from m-sulfobenzoic acid monosodium salt, PG and DMT, optionally but preferably further comprising added PEG, e.g., PEG 3400.

SRA's also include: simple copolymeric blocks of ethylene terephthalate or propylene terephthalate with polyethylene oxide or polypropylene oxide terephthalate, see U.S. 3,959,230 to Hays, May 25, 1976 and U.S. 3,893,929 to Basadur, July 8, 1975; cellulosic derivatives such as the hydroxyether cellulosic polymers available as METHOCEL from Dow; the C₁-C₄ alkyl celluloses and C₄ hydroxyalkyl celluloses, see

U.S. 4,000,093, December 28, 1976 to Nicol. et al.; and the methyl cellulose ethers having an average degree of substitution (methyl) per anhydroglucose unit from about 1.6 to about 2.3 and a solution viscosity of from about 80 to about 120 centipoise measured at 20°C as a 2% aqueous solution. Such materials are available as METOLOSE SM100 and METOLOSE SM200, which are the trade names of methyl cellulose ethers manufactured by Shin-etsu Kagaku Kogyo KK.

Suitable SRA's characterised by poly(vinyl ester) hydrophobe segments include graft copolymers of poly(vinyl ester), e.g., C₁-C₆ vinyl esters, preferably poly(vinyl acetate), grafted onto polyalkylene oxide backbones. See European Patent Application 0 219 048, published April 22, 1987 by Kud, et al. Commercially available examples include SOKALAN SRA's such as SOKALAN HP-22, available from BASF, Germany. Other SRA's are polyesters with repeat units containing 10-15% by weight of ethylene terephthalate together with 80-90% by weight of polyoxyethylene terephthalate derived from a polyoxyethylene glycol of average molecular weight 300-5,000. Commercial examples include ZELCON 5126 from Dupont and MILEASE T from ICI.

Another preferred SRA is an oligomer having empirical formula (CAP)₂(EG/PG)₅(T)₅(SIP)₁ which comprises terephthaloyl (T), sulfoisophthaloyl (SIP), oxyethyleneoxy and oxy-1,2-propylene (EG/PG) units and which is preferably terminated with end-caps (CAP), preferably modified isethionates, as in an oligomer comprising one sulfoisophthaloyl unit, 5 terephthaloyl units, oxyethyleneoxy and oxy-1,2-propyleneoxy units in a defined ratio, preferably about 0.5:1 to about 10:1, and two end-cap units derived from sodium 2-(2-hydroxyethoxy)-ethanesulfonate. Said SRA preferably further comprises from 0.5% to 20%, by weight of the oligomer, of a crystallinity-reducing stabilizer, for example an anionic surfactant such as linear sodium dodecylbenzenesulfonate or a member selected from xylene-, cumene-, and toluene-sulfonates or mixtures thereof, these stabilizers or modifiers being introduced into the synthesis vessel, all as taught in U.S. 5,415,807, Gosselink, Pan, Kellett and Hall, issued May 16, 1995. Suitable monomers for the above SRA include Na-2-(2-hydroxyethoxy)-ethanesulfonate, DMT, Na-dimethyl-5-sulfoisophthalate, EG and PG.

Additional classes of SRA's include: (I) nonionic terephthalates using diisocyanate coupling agents to link polymeric ester structures, see U.S. 4,201,824, Violland et al. and U.S. 4,240,918 Lagasse et al.; and (II) SRA's with carboxylate terminal groups made by adding trimellitic anhydride to known SRA's to convert terminal hydroxyl groups to trimellitate esters. With the proper selection of catalyst, the trimellitic anhydride forms linkages to the terminals of the polymer through an ester of the isolated carboxylic acid of trimellitic anhydride rather than by opening of the

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anhydride linkage. Either nonionic or anionic SRA's may be used as starting materials as long as they have hydroxyl terminal groups which may be esterified. See U.S. 4,525,524 Tung et al.. Other classes include: (III) anionic terephthalate-based SRA's of the urethane-linked variety, see U.S. 4,201,824, Violland et al.; (IV) poly(vinyl caprolactam) and related co-polymers with monomers such as vinyl pyrrolidone and/or dimethylaminoethyl methacrylate, including both nonionic and cationic polymers, see U.S. 4,579,681, Ruppert et al.; (V) graft copolymers, in addition to the SOKALAN types from BASF, made by grafting acrylic monomers onto sulfonated polyesters. These SRA's assertedly have soil release and anti-redeposition activity similar to known cellulose ethers: see EP 279,134 A, 1988, to Rhone-Poulenc Chemie. Still other classes include: (VI) grafts of vinyl monomers such as acrylic acid and vinyl acetate onto proteins such as caseins, see EP 457,205 A to BASF (1991); and (VII) polyesterpolyamide SRA's prepared by condensing adipic acid, caprolactam, and polyethylene glycol, especially for treating polyamide fabrics, see Bevan et al., DE 2,335,044 to Unilever N. V., 1974. Other useful SRA's are described in U.S. Patents 4,240,918, 4,787,989 and 4,525,524.

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Bleaching Compounds - Bleaching Agents and Bleach Activators

The detergent compositions herein may optionally contain bleaching agents or bleaching compositions containing a bleaching agent and one or more bleach activators. When present, bleaching agents will be at levels of from about 0.05% to about 30%, more preferably from about 1% to about 30%, most preferably from about 5% to about 20%, of the detergent composition, especially for fabric laundering. If present, the amount of bleach activators will typically be from about 0.1% to about 60%, more typically from about 0.5% to about 40% of the bleaching composition comprising the bleaching agent-plus-bleach activator.

The bleaching agents used herein can be any of the bleaching agents useful for detergent compositions in textile cleaning that are now known or become known. These include oxygen bleaches as well as other bleaching agents. Perborate bleaches, e.g., sodium perborate (e.g., mono- or tetra-hydrate) can be used herein.

Another category of bleaching agent that can be used without restriction encompasses percarboxylic acid bleaching agents and salts thereof. Suitable examples of this class of agents include magnesium monoperoxyphthalate hexahydrate, the magnesium salt of metachloro perbenzoic acid, 4-nonylamino-4-oxoperoxybutyric acid and diperoxydodecanedioic acid. Such bleaching agents are disclosed in U.S. Patent 4,483,781, Hartman, issued November 20, 1984, U.S. Patent Application 740,446, Burns et al. filed June 3, 1985, European Patent Application 0,133,354, Banks et al, published

February 20, 1985, and U.S. Patent 4,412,934, Chung et al, issued November 1, 1983. Highly preferred bleaching agents also include 6-nonylamino-6-oxoperoxycaproic acid as described in U.S. Patent 4,634,551, issued January 6, 1987 to Burns et al.

Peroxygen bleaching agents can also be used. Suitable peroxygen bleaching compounds include sodium carbonate peroxyhydrate and equivalent "percarbonate" bleaches, sodium pyrophosphate peroxyhydrate, urea peroxyhydrate, and sodium peroxide. Persulfate bleach (e.g., OXONE, manufactured commercially by DuPont) can also be used.

A preferred percarbonate bleach comprises dry particles having an average particle size in the range from about 500 micrometers to about 1,000 micrometers, not more than about 10% by weight of said particles being smaller than about 200 micrometers and not more than about 10% by weight of said particles being larger than about 1,250 micrometers. Optionally, the percarbonate can be coated with silicate, borate or water-soluble surfactants. Percarbonate is available from various commercial sources such as FMC, Solvay and Tokai Denka.

Mixtures of bleaching agents can also be used.

Peroxygen bleaching agents, the perborates, the percarbonates, etc., are preferably combined with bleach activators, which lead to the *in situ* production in aqueous solution (i.e., during the washing process) of the peroxy acid corresponding to the bleach activator. Various nonlimiting examples of activators are disclosed in U.S. Patent 4,915,854, issued April 10, 1990 to Mao et al, and U.S. Patent 4,412,934. The nonanoyloxybenzene sulfonate (NOBS) and tetraacetyl ethylene diamine (TAED) activators are typical, and mixtures thereof can also be used. See also U.S. 4,634,551 for other typical bleaches and activators useful herein.

Highly preferred amido-derived bleach activators are those of the formulae: $R^1N(R^5)C(O)R^2C(O)L$ or $R^1C(O)N(R^5)R^2C(O)L$ wherein R^1 is an alkyl group containing from about 6 to about 12 carbon atoms, R^2 is an alkylene containing from 1 to about 6 carbon atoms, R^5 is H or alkyl, aryl, or alkaryl containing from about 1 to about 10 carbon atoms, and L is any suitable leaving group. A leaving group is any group that is displaced from the bleach activator as a consequence of the nucleophilic attack on the bleach activator by the perhydrolysis anion. A preferred leaving group is phenyl sulfonate.

Preferred examples of bleach activators of the above formulae include (6-octanamido-caproyl)oxybenzenesulfonate, (6-nonanamidocaproyl)oxybenzenesulfonate, (6-decanamido-caproyl)oxybenzenesulfonate, and mixtures thereof as described in U.S. Patent 4,634,551, incorporated herein by reference.

Another class of bleach activators comprises the benzoxazin-type activators disclosed by Hodge et al in U.S. Patent 4,966,723, issued October 30, 1990, incorporated herein by reference. A highly preferred activator of the benzoxazin-type is:

Still another class of preferred bleach activators includes the acyl lactam activators, especially acyl caprolactams and acyl valerolactams of the formulae:

wherein R⁶ is H or an alkyl, aryl, alkoxyaryl, or alkaryl group containing from 1 to about 12 carbon atoms. Highly preferred lactam activators include benzoyl caprolactam, octanoyl caprolactam, 3,5,5-trimethylhexanoyl caprolactam, nonanoyl caprolactam, decanoyl caprolactam, undecenoyl caprolactam, benzoyl valerolactam, octanoyl valerolactam, decanoyl valerolactam, undecenoyl valerolactam, nonanoyl valerolactam, 3,5,5-trimethylhexanoyl valerolactam and mixtures thereof. See also U.S. Patent 4,545,784, issued to Sanderson, October 8, 1985, incorporated herein by reference, which discloses acyl caprolactams, including benzoyl caprolactam, adsorbed into sodium perborate.

Bleaching agents other than oxygen bleaching agents are also known in the art and can be utilized herein. One type of non-oxygen bleaching agent of particular interest includes photoactivated bleaching agents such as the sulfonated zinc and/or aluminum phthalocyanines. See U.S. Patent 4,033,718, issued July 5, 1977 to Holcombe et al. If used, detergent compositions will typically contain from about 0.025% to about 1.25%, by weight, of such bleaches, especially sulfonate zinc phthalocyanine.

If desired, the bleaching compounds can be catalyzed by means of a manganese compound. Such compounds are well known in the art and include, for example, the manganese-based catalysts disclosed in U.S. Pat. 5,246,621, U.S. Pat. 5,244,594; U.S. Pat. 5,194,416; U.S. Pat. 5,114,606; and European Pat. App. Pub. Nos. 549,271A1, 549,272A1, 544,440A2, and 544,490A1; Preferred examples of these catalysts include Mn^{IV}₂(u-O)₃(1,4,7-trimethyl-1,4,7-triazacyclononane)₂(PF₆)₂, Mn^{III}₂(u-O)₁(u-OAc)₂(1,4,7-trimethyl-1,4,7-triazacyclononane)₂.(ClO₄)₂, Mn^{IV}₄(u-O)₆(1,4,7-trimethyl-1,4,7-triazacyclononane)₂.(ClO₄)₂, Mn^{IV}₄(u-O)₆(1,4,7-trimethyl-1,4,7-triazacyclononane)₂.(ClO₄)₂, Mn^{IV}₄(u-O)₆(1,4,7-trimethyl-1,4,7-triazacyclononane)₂.

triazacyclononane)₄(ClO₄)₄, Mn^{III}Mn^{IV}₄(u-O)₁(u-OAc)₂-(1,4,7-trimethyl-1,4,7-triazacyclononane)₂(ClO₄)₃, Mn^{IV}(1,4,7-trimethyl-1,4,7-triazacyclononane)-(OCH₃)₃(PF₆), and mixtures thereof. Other metal-based bleach catalysts include those disclosed in U.S. Pat. 4,430,243 and U.S. Pat. 5,114,611. The use of manganese with various complex ligands to enhance bleaching is also reported in the following United States Patents: 4,728,455; 5,284,944; 5,246,612; 5,256,779; 5,280,117; 5,274,147; 5,153,161; and 5,227,084.

As a practical matter, and not by way of limitation, the compositions and processes herein can be adjusted to provide on the order of at least one part per ten million of the active bleach catalyst species in the aqueous washing liquor, and will preferably provide from about 0.1 ppm to about 700 ppm, more preferably from about 1 ppm to about 500 ppm, of the catalyst species in the laundry liquor.

A wide variety of other ingredients useful in detergent compositions can be included in the compositions herein, including other active ingredients, carriers, hydrotropes, processing aids, dyes or pigments, solvents for liquid formulations, solid fillers for bar compositions, etc. If high sudsing is desired, suds boosters such as the C₁₀-C₁₆ alkanolamides can be incorporated into the compositions, typically at 1%-10% levels. The C₁₀-C₁₄ monoethanol and diethanol amides illustrate a typical class of such suds boosters. Use of such suds boosters with high sudsing adjunct surfactants such as the amine oxides, betaines and sultaines noted above is also advantageous. If desired, soluble magnesium salts such as MgCl₂, MgSO₄, and the like, can be added at levels of, typically, 0.1%-2%, to provide additional suds and to enhance grease removal performance.

Various detersive ingredients employed in the present compositions optionally can be further stabilized by absorbing said ingredients onto a porous hydrophobic substrate, then coating said substrate with a hydrophobic coating. Preferably, the detersive ingredient is admixed with a surfactant before being absorbed into the porous substrate. In use, the detersive ingredient is released from the substrate into the aqueous washing liquor, where it performs its intended detersive function.

To illustrate this technique in more detail, a porous hydrophobic silica (trademark SIPERNAT D10, DeGussa) is admixed with a proteolytic enzyme solution containing 3%-5% of C₁₃₋₁₅ ethoxylated alcohol (EO 7) nonionic surfactant. Typically, the enzyme/surfactant solution is 2.5 X the weight of silica. The resulting powder is dispersed with stirring in silicone oil (various silicone oil viscosities in the range of 500-12,500 can be used). The resulting silicone oil dispersion is emulsified or otherwise added to the final detergent matrix. By this means, ingredients such as the

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aforementioned enzymes, bleaches, bleach activators, bleach catalysts, photoactivators, dyes, fluorescers, fabric conditioners and hydrolyzable surfactants can be "protected" for use in detergents, including liquid laundry detergent compositions.

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Liquid detergent compositions can contain water and other solvents as carriers. Low molecular weight primary or secondary alcohols exemplified by methanol, ethanol, propanol, and isopropanol are suitable. Monohydric alcohols are preferred for solubilizing surfactant, but polyols such as those containing from 2 to about 6 carbon atoms and from 2 to about 6 hydroxy groups (e.g., 1,3-propanediol, ethylene glycol, glycerin, and 1,2-propanediol) can also be used. The compositions may contain from 5% to 90%, typically 10% to 50% of such carriers.

The detergent compositions herein will preferably be formulated such that, during use in aqueous cleaning operations, the wash water will have a pH of between about 6.5 and about 11, preferably between about 7.5 and 10.5. Laundry products are typically at pH 9-11. Techniques for controlling pH at recommended usage levels include the use of buffers, alkalis, acids, etc., and are well known to those skilled in the art.

Enzymes

Enzymes can be included in the present detergent compositions for a variety of purposes, including removal of protein-based, carbohydrate-based, or triglyceride-based stains from surfaces such as textiles, for the prevention of refugee dye transfer, for example in laundering, and for fabric restoration. Suitable enzymes include proteases, amylases, lipases, cellulases, peroxidases, and mixtures thereof of any suitable origin, such as vegetable, animal, bacterial, fungal and yeast origin. Preferred selections are influenced by factors such as pH-activity and/or stability optima, thermostability, and stability to active detergents, builders and the like. In this respect bacterial or fungal enzymes are preferred, such as bacterial amylases and proteases, and fungal cellulases.

"Detersive enzyme", as used herein, means any enzyme having a cleaning, stain removing or otherwise beneficial effect in a laundry, hard surface cleaning or personal care detergent composition. Preferred detersive enzymes are hydrolases such as proteases, amylases and lipases. Preferred enzymes for laundry purposes include, but are not limited to, proteases, cellulases, lipases and peroxidases.

Enzymes are normally incorporated into detergent or detergent additive compositions at levels sufficient to provide a "cleaning-effective amount". The term "cleaning effective amount" refers to any amount capable of producing a cleaning, stain removal, soil removal, whitening, deodorizing, or freshness improving effect on substrates such as fabrics. In practical terms for current commercial preparations, typical amounts are up to about 5 mg by weight, more typically 0.01 mg to 3 mg, of active

enzyme per gram of the detergent composition. Stated otherwise, the compositions herein will typically comprise from 0.001% to 5%, preferably 0.01%-1% by weight of a commercial enzyme preparation. Protease enzymes are usually present in such commercial preparations at levels sufficient to provide from 0.005 to 0.1 Anson units (AU) of activity per gram of composition. For certain detergents, it may be desirable to increase the active enzyme content of the commercial preparation in order to minimize the total amount of non-catalytically active materials and thereby improve spotting/filming or other end-results. Higher active levels may also be desirable in highly concentrated detergent formulations.

Suitable examples of proteases are the subtilisins which are obtained from particular strains of B. subtilis and B. licheniformis. One suitable protease is obtained from a strain of Bacillus, having maximum activity throughout the pH range of 8-12, developed and sold as ESPERASE® by Novo Industries A/S of Denmark, hereinafter "Novo". The preparation of this enzyme and analogous enzymes is described in GB 1,243,784 to Novo. Other suitable proteases include ALCALASE® and SAVINASE® from Novo and MAXATASE® from International Bio-Synthetics, Inc., The Netherlands; as well as Protease A as disclosed in EP 130,756 A, January 9, 1985 and Protease B as disclosed in EP 303,761 A, April 28, 1987 and EP 130,756 A, January 9, 1985. See also a high pH protease from Bacillus sp. NCIMB 40338 described in WO 9318140 A to Novo. Enzymatic detergents comprising protease, one or more other enzymes, and a reversible protease inhibitor are described in WO 9203529 A to Novo. Other preferred proteases include those of WO 9510591 A to Procter & Gamble . When desired, a protease having decreased adsorption and increased hydrolysis is available as described in WO 9507791 to Procter & Gamble. A recombinant trypsin-like protease for detergents suitable herein is described in WO 9425583 to Novo.

The preferred liquid laundry detergent compositions according to the present invention further comprise at least 0.001% by weight, of a protease enzyme. However, an effective amount of protease enzyme is sufficient for use in the liquid laundry detergent compositions described herein. The term "an effective amount" refers to any amount capable of producing a cleaning, stain removal, soil removal, whitening, deodorizing, or freshness improving effect on substrates such as fabrics. In practical terms for current commercial preparations, typical amounts are up to about 5 mg by weight, more typically 0.01 mg to 3 mg, of active enzyme per gram of the detergent composition. Stated otherwise, the compositions herein will typically comprise from 0.001% to 5%, preferably 0.01%-1% by weight of a commercial enzyme preparation. The protease enzymes of the present invention are usually present in such commercial

preparations at levels sufficient to provide from 0.005 to 0.1 Anson units (AU) of activity per gram of composition.

Preferred liquid laundry detergent compositions of the present invention comprise a protease enzyme, referred to as "Protease D", which is a carbonyl hydrolase variant having an amino acid sequence not found in nature, which is derived from a precursor carbonyl hydrolase by substituting a different amino acid for a plurality of amino acid residues at a position in said carbonyl hydrolase equivalent to position +76, preferably also in combination with one or more amino acid residue positions equivalent to those selected from the group consisting of +99, +101, +103, +104, +107, +123, +27, +105, +109, +126, +128, +135, +156, +166, +195, +197, +204, +206, +210, +216, +217, +218, +222, +260, +265, and/or +274 according to the numbering of *Bacillus amyloliquefaciens* subtilisin, as described in WO 95/10615 published April 20, 1995 by Genencor International.

Useful proteases are also described in PCT publications: WO 95/30010 published November 9, 1995 by The Procter & Gamble Company; WO 95/30011 published November 9, 1995 by The Procter & Gamble Company; WO 95/29979 published November 9, 1995 by The Procter & Gamble Company.

Preferred proteolytic enzymes are also modified bacterial serine proteases, such as those described in European Patent Application Serial Number 87 303,761.8, filed April 28, 1987 (particularly pages 17, 24 and 98), and which is called herein "Protease B", and in European Patent Application 199,404, Venegas, published October 29, 1986, which refers to a modified bacterial serine proteolytic enzyme which is called "Protease A" herein, Protease A as disclosed in EP 130,756 A, January 9, 1985 and Protease B as disclosed in EP 303,761 A, April 28, 1987 and EP 130,756 A, January 9, 1985

Amylases suitable herein, include, for example, α-amylases described in GB 1,296,839 to Novo; RAPIDASE®, International Bio-Synthetics, Inc. and TERMAMYL®, Novo. FUNGAMYL® from Novo is especially useful. Engineering of enzymes for improved stability, e.g., oxidative stability, is known. See, for example J. Biological Chem., Vol. 260, No. 11, June 1985, pp 6518-6521. Certain preferred embodiments of the present compositions can make use of amylases having improved stability in detergents, especially improved oxidative stability as measured against a reference-point of TERMAMYL® in commercial use in 1993. These preferred amylases herein share the characteristic of being "stability-enhanced" amylases, characterized, at a minimum, by a measurable improvement in one or more of: oxidative stability, e.g., to hydrogen peroxide / tetraacetylethylene-diamine in buffered solution at pH 9-10; thermal stability, e.g., at common wash temperatures such as about 60°C; or alkaline stability,

e.g., at a pH from about 8 to about 11, measured versus the above-identified referencepoint amylase. Stability can be measured using any of the art-disclosed technical tests. See, for example, references disclosed in WO 9402597. Stability-enhanced amylases can be obtained from Novo or from Genencor International. One class of highly preferred amylases herein have the commonality of being derived using site-directed mutagenesis from one or more of the *Baccillus* amylases, especially the *Bacillus* α-amylases. regardless of whether one, two or multiple amylase strains are the immediate precursors. Oxidative stability-enhanced amylases vs. the above-identified reference amylase are preferred for use, especially in bleaching, more preferably oxygen bleaching, as distinct from chlorine bleaching, detergent compositions herein. Such preferred amylases include (a) an amylase according to the hereinbefore incorporated WO 9402597, Novo, Feb. 3, 1994, as further illustrated by a mutant in which substitution is made, using alanine or threonine, preferably threonine, of the methionine residue located in position 197 of the B.licheniformis alpha-amylase, known as TERMAMYL®, or the homologous position variation of a similar parent amylase, such as B. amyloliquefaciens, B. subtilis, or B. stearothermophilus; (b) stability-enhanced amylases as described by Genencor International in a paper entitled "Oxidatively Resistant alpha-Amylases" presented at the 207th American Chemical Society National Meeting, March 13-17 1994, by C. Mitchinson. Therein it was noted that bleaches in automatic dishwashing detergents inactivate alpha-amylases but that improved oxidative stability amylases have been made by Genencor from B.licheniformis NCIB8061. Methionine (Met) was identified as the most likely residue to be modified. Met was substituted, one at a time, in positions 8, 15, 197, 256, 304, 366 and 438 leading to specific mutants, particularly important being M197L and M197T with the M197T variant being the most stable expressed variant. Stability was measured in CASCADE® and SUNLIGHT®; (c) particularly preferred amylases herein include amylase variants having additional modification in the immediate parent as described in WO 9510603 A and are available from the assignee, Novo, as DURAMYL®. Other particularly preferred oxidative stability enhanced amylase include those described in WO 9418314 to Genencor International and WO 9402597 to Novo. Any other oxidative stability-enhanced amylase can be used, for example as derived by site-directed mutagenesis from known chimeric, hybrid or simple mutant parent forms of available amylases. Other preferred enzyme modifications are accessible. See WO 9509909 A to Novo.

Cellulases usable herein include both bacterial and fungal types, preferably having a pH optimum between 5 and 9.5. U.S. 4,435,307, Barbesgoard et al, March 6, 1984, discloses suitable fungal cellulases from *Humicola insolens* or *Humicola* strain

DSM1800 or a cellulase 212-producing fungus belonging to the genus *Aeromonas*, and cellulase extracted from the hepatopancreas of a marine mollusk, *Dolabella Auricula Solander*. Suitable cellulases are also disclosed in GB-A-2.075.028; GB-A-2.095.275 and DE-OS-2.247.832. CAREZYME® (Novo) is especially useful. See also WO 9117243 to Novo.

Suitable lipase enzymes for detergent usage include those produced by microorganisms of the *Pseudomonas* group, such as *Pseudomonas stutzeri* ATCC 19.154, as disclosed in GB 1,372,034. See also lipases in Japanese Patent Application 53,20487, laid open Feb. 24, 1978. This lipase is available from Amano Pharmaceutical Co. Ltd., Nagoya, Japan, under the trade name Lipase P "Amano," or "Amano-P." Other suitable commercial lipases include Amano-CES, lipases ex *Chromobacter viscosum*, e.g. *Chromobacter viscosum var. lipolyticum* NRRLB 3673 from Toyo Jozo Co., Tagata, Japan; *Chromobacter viscosum* lipases from U.S. Biochemical Corp., U.S.A. and Disoynth Co., The Netherlands, and lipases ex *Pseudomonas gladioli*. LIPOLASE® enzyme derived from *Humicola lanuginosa* and commercially available from Novo, see also EP 341,947, is a preferred lipase for use herein. Lipase and amylase variants stabilized against peroxidase enzymes are described in WO 9414951 A to Novo. See also WO 9205249 and RD 94359044.

Cutinase enzymes suitable for use herein are described in WO 8809367 A to Genencor.

Peroxidase enzymes may be used in combination with oxygen sources, e.g., percarbonate, perborate, hydrogen peroxide, etc., for "solution bleaching" or prevention of transfer of dyes or pigments removed from substrates during the wash to other substrates present in the wash solution. Known peroxidases include horseradish peroxidase, ligninase, and haloperoxidases such as chloro- or bromo-peroxidase. Peroxidase-containing detergent compositions are disclosed in WO 89099813 A, October 19, 1989 to Novo and WO 8909813 A to Novo.

A range of enzyme materials and means for their incorporation into synthetic detergent compositions is also disclosed in WO 9307263 A and WO 9307260 A to Genencor International, WO 8908694 A to Novo, and U.S. 3,553,139, January 5, 1971 to McCarty et al. Enzymes are further disclosed in U.S. 4,101,457, Place et al, July 18, 1978, and in U.S. 4,507,219, Hughes, March 26, 1985. Enzyme materials useful for liquid detergent formulations, and their incorporation into such formulations, are disclosed in U.S. 4,261,868, Hora et al, April 14, 1981. Enzymes for use in detergents can be stabilized by various techniques. Enzyme stabilization techniques are disclosed and exemplified in U.S. 3,600,319, August 17, 1971, Gedge et al, EP 199,405 and EP

200,586, October 29, 1986, Venegas. Enzyme stabilization systems are also described, for example, in U.S. 3,519,570. A useful Bacillus, sp. AC13 giving proteases, xylanases and cellulases, is described in WO 9401532 A to Novo.

Enzyme Stabilizing System

Enzyme-containing, including but not limited to, liquid compositions, herein may comprise from about 0.001% to about 10%, preferably from about 0.005% to about 8%, most preferably from about 0.01% to about 6%, by weight of an enzyme stabilizing system. The enzyme stabilizing system can be any stabilizing system which is compatible with the detersive enzyme. Such a system may be inherently provided by other formulation actives, or be added separately, e.g., by the formulator or by a manufacturer of detergent-ready enzymes. Such stabilizing systems can, for example, comprise calcium ion, boric acid, propylene glycol, short chain carboxylic acids, boronic acids, and mixtures thereof, and are designed to address different stabilization problems depending on the type and physical form of the detergent composition.

One stabilizing approach is the use of water-soluble sources of calcium and/or magnesium ions in the finished compositions which provide such ions to the enzymes. Calcium ions are generally more effective than magnesium ions and are preferred herein if only one type of cation is being used. Typical detergent compositions, especially liquids, will comprise from about 1 to about 30, preferably from about 2 to about 20, more preferably from about 8 to about 12 millimoles of calcium ion per liter of finished detergent composition, though variation is possible depending on factors including the multiplicity, type and levels of enzymes incorporated. Preferably water-soluble calcium or magnesium salts are employed, including for example calcium chloride, calcium hydroxide, calcium formate, calcium malate, calcium maleate, calcium hydroxide and calcium acetate; more generally, calcium sulfate or magnesium salts corresponding to the exemplified calcium salts may be used. Further increased levels of Calcium and/or Magnesium may of course be useful, for example for promoting the grease-cutting action of certain types of surfactant.

Another stabilizing approach is by use of borate species. See Severson, U.S. 4,537,706. Borate stabilizers, when used, may be at levels of up to 10% or more of the composition though more typically, levels of up to about 3% by weight of boric acid or other borate compounds such as borax or orthoborate are suitable for liquid detergent use. Substituted boric acids such as phenylboronic acid, butaneboronic acid, p-bromophenylboronic acid or the like can be used in place of boric acid and reduced levels of total boron in detergent compositions may be possible though the use of such substituted boron derivatives.

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Stabilizing systems of certain cleaning compositions may further comprise from 0 to about 10%, preferably from about 0.01% to about 6% by weight, of chlorine bleach scavengers, added to prevent chlorine bleach species present in many water supplies from attacking and inactivating the enzymes, especially under alkaline conditions. While chlorine levels in water may be small, typically in the range from about 0.5 ppm to about 1.75 ppm, the available chlorine in the total volume of water that comes in contact with the enzyme, for example during fabric-washing, can be relatively large; accordingly, enzyme stability to chlorine in-use is sometimes problematic. Since perborate or percarbonate, which have the ability to react with chlorine bleach, may present in certain of the instant compositions in amounts accounted for separately from the stabilizing system, the use of additional stabilizers against chlorine, may, most generally, not be essential, though improved results may be obtainable from their use. Suitable chlorine scavenger anions are widely known and readily available, and, if used, can be salts containing ammonium cations with sulfite, bisulfite, thiosulfite, thiosulfate, iodide, etc. Antioxidants such as carbamate, ascorbate, etc., organic amines such as ethylenediaminetetracetic acid (EDTA) or alkali metal salt thereof, monoethanolamine (MEA), and mixtures thereof can likewise be used. Likewise, special enzyme inhibition systems can be incorporated such that different enzymes have maximum compatibility. Other conventional scavengers such as bisulfate, nitrate, chloride, sources of hydrogen peroxide such as sodium perborate tetrahydrate, sodium perborate monohydrate and sodium percarbonate, as well as phosphate, condensed phosphate, acetate, benzoate, citrate, formate, lactate, malate, tartrate, salicylate, etc., and mixtures thereof can be used if desired. In general, since the chlorine scavenger function can be performed by ingredients separately listed under better recognized functions, (e.g., hydrogen peroxide sources), there is no absolute requirement to add a separate chlorine scavenger unless a compound performing that function to the desired extent is absent from an enzymecontaining embodiment of the invention; even then, the scavenger is added only for optimum results. Moreover, the formulator will exercise a chemist's normal skill in avoiding the use of any enzyme scavenger or stabilizer which is majorly incompatible, as formulated, with other reactive ingredients, if used. In relation to the use of ammonium salts, such salts can be simply admixed with the detergent composition but are prone to adsorb water and/or liberate ammonia during storage. Accordingly, such materials, if present, are desirably protected in a particle such as that described in US 4,652,392, Baginski et al.

Builders

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Detergent builders can optionally be included in the compositions herein to assist in controlling mineral hardness. Inorganic as well as organic builders can be used. Builders are typically used in fabric laundering compositions to assist in the removal of particulate soils.

The level of builder can vary widely depending upon the end use of the composition and its desired physical form. When present, the compositions will typically comprise at least about 1% builder. Liquid formulations typically comprise from about 5% to about 50%, more typically about 5% to about 30%, by weight, of detergent builder. Granular formulations typically comprise from about 10% to about 80%, more typically from about 15% to about 50% by weight, of the detergent builder. Lower or higher levels of builder, however, are not meant to be excluded.

Inorganic or P-containing detergent builders include, but are not limited to, the alkali metal, ammonium and alkanolammonium salts of polyphosphates (exemplified by the tripolyphosphates, pyrophosphates, and glassy polymeric meta-phosphates), phosphonates, phytic acid, silicates, carbonates (including bicarbonates and sesquicarbonates), sulphates, and aluminosilicates. However, non-phosphate builders are required in some locales. Importantly, the compositions herein function surprisingly well even in the presence of the so-called "weak" builders (as compared with phosphates) such as citrate, or in the so-called "underbuilt" situation that may occur with zeolite or layered silicate builders.

Examples of silicate builders are the alkali metal silicates, particularly those having a SiO₂:Na₂O ratio in the range 1.6:1 to 3.2:1 and layered silicates, such as the layered sodium silicates described in U.S. Patent 4,664,839, issued May 12, 1987 to H. P. Rieck. NaSKS-6 is the trademark for a crystalline layered silicate marketed by Hoechst (commonly abbreviated herein as "SKS-6"). Unlike zeolite builders, the Na SKS-6 silicate builder does not contain aluminum. NaSKS-6 has the delta-Na₂SiO₅ morphology form of layered silicate. It can be prepared by methods such as those described in German DE-A-3,417,649 and DE-A-3,742,043. SKS-6 is a highly preferred layered silicate for use herein, but other such layered silicates, such as those having the general formula NaMSi_xO_{2x+1}·yH₂O wherein M is sodium or hydrogen, x is a number from 1.9 to 4, preferably 2, and y is a number from 0 to 20, preferably 0 can be used herein. Various other layered silicates from Hoechst include NaSKS-5, NaSKS-7 and NaSKS-11, as the alpha, beta and gamma forms. As noted above, the delta-Na₂SiO₅ (NaSKS-6 form) is most preferred for use herein. Other silicates may also be useful such as for example magnesium silicate, which can serve as a crispening agent in granular

formulations, as a stabilizing agent for oxygen bleaches, and as a component of suds control systems.

Examples of carbonate builders are the alkaline earth and alkali metal carbonates as disclosed in German Patent Application No. 2,321,001 published on November 15, 1973.

Aluminosilicate builders are useful in the present invention. Aluminosilicate builders are of great importance in most currently marketed heavy duty granular detergent compositions, and can also be a significant builder ingredient in liquid detergent formulations. Aluminosilicate builders include those having the empirical formula:

$M_z(zAlO_2)_y$ }·xH₂O

wherein z and y are integers of at least 6, the molar ratio of z to y is in the range from 1.0 to about 0.5, and x is an integer from about 15 to about 264.

Useful aluminosilicate ion exchange materials are commercially available. These aluminosilicates can be crystalline or amorphous in structure and can be naturally-occurring aluminosilicates or synthetically derived. A method for producing aluminosilicate ion exchange materials is disclosed in U.S. Patent 3,985,669, Krummel, et al, issued October 12, 1976. Preferred synthetic crystalline aluminosilicate ion exchange materials useful herein are available under the designations Zeolite A, Zeolite P (B), Zeolite MAP and Zeolite X. In an especially preferred embodiment, the crystalline aluminosilicate ion exchange material has the formula:

$$Na_{12}[(AIO_2)_{12}(SiO_2)_{12}] \cdot xH_2O$$

wherein x is from about 20 to about 30, especially about 27. This material is known as Zeolite A. Dehydrated zeolites (x = 0 - 10) may also be used herein. Preferably, the aluminosilicate has a particle size of about 0.1-10 microns in diameter.

Organic detergent builders suitable for the purposes of the present invention include, but are not restricted to, a wide variety of polycarboxylate compounds. As used herein, "polycarboxylate" refers to compounds having a plurality of carboxylate groups, preferably at least 3 carboxylates. Polycarboxylate builder can generally be added to the composition in acid form, but can also be added in the form of a neutralized salt. When utilized in salt form, alkali metals, such as sodium, potassium, and lithium, or alkanolammonium salts are preferred.

Included among the polycarboxylate builders are a variety of categories of useful materials. One important category of polycarboxylate builders encompasses the ether polycarboxylates, including oxydisuccinate, as disclosed in Berg, U.S. Patent 3,128,287, issued April 7, 1964, and Lamberti et al, U.S. Patent 3,635,830, issued January 18, 1972.

See also "TMS/TDS" builders of U.S. Patent 4,663,071, issued to Bush et al, on May 5, 1987. Suitable ether polycarboxylates also include cyclic compounds, particularly alicyclic compounds, such as those described in U.S. Patents 3,923,679; 3,835,163; 4,158,635; 4,120,874 and 4,102,903.

Other useful detergency builders include the ether hydroxypolycarboxylates, copolymers of maleic anhydride with ethylene or vinyl methyl ether, 1, 3, 5-trihydroxy benzene-2, 4, 6-trisulphonic acid, and carboxymethyloxysuccinic acid, the various alkali metal, ammonium and substituted ammonium salts of polyacetic acids such as ethylenediamine tetraacetic acid and nitrilotriacetic acid, as well as polycarboxylates such as mellitic acid, succinic acid, oxydisuccinic acid, polymaleic acid, benzene 1,3,5-tricarboxylic acid, carboxymethyloxysuccinic acid, and soluble salts thereof.

Citrate builders, e.g., citric acid and soluble salts thereof (particularly sodium salt), are polycarboxylate builders of particular importance for heavy duty liquid detergent formulations due to their availability from renewable resources and their biodegradability. Citrates can also be used in granular compositions, especially in combination with zeolite and/or layered silicate builders. Oxydisuccinates are also especially useful in such compositions and combinations.

Also suitable in the detergent compositions of the present invention are the 3,3-dicarboxy-4-oxa-1,6-hexanedioates and the related compounds disclosed in U.S. Patent 4,566,984, Bush, issued January 28, 1986. Useful succinic acid builders include the C5-C20 alkyl and alkenyl succinic acids and salts thereof. A particularly preferred compound of this type is dodecenylsuccinic acid. Specific examples of succinate builders include: laurylsuccinate, myristylsuccinate, palmitylsuccinate, 2-dodecenylsuccinate (preferred), 2-pentadecenylsuccinate, and the like. Laurylsuccinates are the preferred builders of this group, and are described in European Patent Application 86200690.5/0,200,263, published November 5, 1986.

Other suitable polycarboxylates are disclosed in U.S. Patent 4,144,226, Crutchfield et al, issued March 13, 1979 and in U.S. Patent 3,308,067, Diehl, issued March 7, 1967. See also Diehl U.S. Patent 3,723,322.

Fatty acids, e.g., C₁₂-C₁₈ monocarboxylic acids, can also be incorporated into the compositions alone, or in combination with the aforesaid builders, especially citrate and/or the succinate builders, to provide additional builder activity. Such use of fatty acids will generally result in a diminution of sudsing, which should be taken into account by the formulator.

In situations where phosphorus-based builders can be used, and especially in the formulation of bars used for hand-laundering operations, the various alkali metal

phosphates such as the well-known sodium tripolyphosphates, sodium pyrophosphate and sodium orthophosphate can be used. Phosphonate builders such as ethane-1-hydroxy-1,1-diphosphonate and other known phosphonates (see, for example, U.S. Patents 3,159,581; 3,213,030; 3,422,021; 3,400,148 and 3,422,137) can also be used.

Chelating Agents

The detergent compositions herein may also optionally contain one or more iron and/or manganese chelating agents. Such chelating agents can be selected from the group consisting of amino carboxylates, amino phosphonates, polyfunctionally-substituted aromatic chelating agents and mixtures therein, all as hereinafter defined. Without intending to be bound by theory, it is believed that the benefit of these materials is due in part to their exceptional ability to remove iron and manganese ions from washing solutions by formation of soluble chelates.

Amino carboxylates useful as optional chelating agents include ethylenediaminetetracetates, N-hydroxyethylethylenediaminetriacetates, nitrilotriacetates, ethylenediamine tetraproprionates, triethylenetetraaminehexacetates, diethylenetriaminepentaacetates, and ethanoldiglycines, alkali metal, ammonium, and substituted ammonium salts therein and mixtures therein.

Amino phosphonates are also suitable for use as chelating agents in the compositions of the invention when at lease low levels of total phosphorus are permitted in detergent compositions, and include ethylenediaminetetrakis (methylenephosphonates) as DEQUEST. Preferred, these amino phosphonates to not contain alkyl or alkenyl groups with more than about 6 carbon atoms.

Polyfunctionally-substituted aromatic chelating agents are also useful in the compositions herein. See U.S. Patent 3,812,044, issued May 21, 1974, to Connor et al. Preferred compounds of this type in acid form are dihydroxydisulfobenzenes such as 1,2-dihydroxy-3,5-disulfobenzene.

A preferred biodegradable chelator for use herein is ethylenediamine disuccinate ("EDDS"), especially the [S,S] isomer as described in U.S. Patent 4,704,233, November 3, 1987, to Hartman and Perkins.

If utilized, these chelating agents will generally comprise from about 0.1% to about 10% by weight of the detergent compositions herein. More preferably, if utilized, the chelating agents will comprise from about 0.1% to about 3.0% by weight of such compositions.

Clay Soil Removal/Anti-redeposition Agents

The compositions of the present invention can also optionally contain watersoluble ethoxylated amines having clay soil removal and antiredeposition properties. Granular detergent compositions which contain these compounds typically contain from about 0.01% to about 10.0% by weight of the water-soluble ethoxylates amines; liquid detergent compositions typically contain about 0.01% to about 5%.

The most preferred soil release and anti-redeposition agent is ethoxylated tetraethylenepentamine. Exemplary ethoxylated amines are further described in U.S. Patent 4,597,898, VanderMeer, issued July 1, 1986. Another group of preferred clay soil removal-antiredeposition agents are the cationic compounds disclosed in European Patent Application 111,965, Oh and Gosselink, published June 27, 1984. Other clay soil removal/antiredeposition agents which can be used include the ethoxylated amine polymers disclosed in European Patent Application 111,984, Gosselink, published June 27, 1984; the zwitterionic polymers disclosed in European Patent Application 112,592, Gosselink, published July 4, 1984; and the amine oxides disclosed in U.S. Patent 4,548,744, Connor, issued October 22, 1985. Other clay soil removal and/or anti redeposition agents known in the art can also be utilized in the compositions herein. Another type of preferred antiredeposition agent includes the carboxy methyl cellulose (CMC) materials. These materials are well known in the art.

Polymeric Dispersing Agents

Polymeric dispersing agents can advantageously be utilized at levels from about 0.1% to about 7%, by weight, in the compositions herein, especially in the presence of zeolite and/or layered silicate builders. Suitable polymeric dispersing agents include polymeric polycarboxylates and polyethylene glycols, although others known in the art can also be used. It is believed, though it is not intended to be limited by theory, that polymeric dispersing agents enhance overall detergent builder performance, when used in combination with other builders (including lower molecular weight polycarboxylates) by crystal growth inhibition, particulate soil release peptization, and anti-redeposition.

Polymeric polycarboxylate materials can be prepared by polymerizing or copolymerizing suitable unsaturated monomers, preferably in their acid form.

Unsaturated monomeric acids that can be polymerized to form suitable polymeric polycarboxylates include acrylic acid, maleic acid (or maleic anhydride), fumaric acid, itaconic acid, aconitic acid, mesaconic acid, citraconic acid and methylenemalonic acid. The presence in the polymeric polycarboxylates herein or monomeric segments, containing no carboxylate radicals such as vinylmethyl ether, styrene, ethylene, etc. is suitable provided that such segments do not constitute more than about 40% by weight.

Particularly suitable polymeric polycarboxylates can be derived from acrylic acid. Such acrylic acid-based polymers which are useful herein are the water-soluble salts of polymerized acrylic acid. The average molecular weight of such polymers in the acid

form preferably ranges from about 2,000 to 10,000, more preferably from about 4,000 to 7,000 and most preferably from about 4,000 to 5,000. Water-soluble salts of such acrylic acid polymers can include, for example, the alkali metal, ammonium and substituted ammonium salts. Soluble polymers of this type are known materials. Use of polyacrylates of this type in detergent compositions has been disclosed, for example, in Diehl, U.S. Patent 3,308,067, issued march 7, 1967.

Acrylic/maleic-based copolymers may also be used as a preferred component of the dispersing/anti-redeposition agent. Such materials include the water-soluble salts of copolymers of acrylic acid and maleic acid. The average molecular weight of such copolymers in the acid form preferably ranges from about 2,000 to 100,000, more preferably from about 5,000 to 75,000, most preferably from about 7,000 to 65,000. The ratio of acrylate to maleate segments in such copolymers will generally range from about 30:1 to about 1:1, more preferably from about 10:1 to 2:1. Water-soluble salts of such acrylic acid/maleic acid copolymers can include, for example, the alkali metal, ammonium and substituted ammonium salts. Soluble acrylate/maleate copolymers of this type are known materials which are described in European Patent Application No. 66915, published December 15, 1982, as well as in EP 193,360, published September 3, 1986, which also describes such polymers comprising hydroxypropylacrylate. Still other useful dispersing agents include the maleic/acrylic/vinyl alcohol terpolymers. Such materials are also disclosed in EP 193,360, including, for example, the 45/45/10 terpolymer of acrylic/maleic/vinyl alcohol.

Another polymeric material which can be included is polyethylene glycol (PEG). PEG can exhibit dispersing agent performance as well as act as a clay soil removalantiredeposition agent. Typical molecular weight ranges for these purposes range from about 500 to about 100,000, preferably from about 1,000 to about 50,000, more preferably from about 1,500 to about 10,000.

Polyaspartate and polyglutamate dispersing agents may also be used, especially in conjunction with zeolite builders. Dispersing agents such as polyaspartate preferably have a molecular weight (avg.) of about 10,000.

Brightener

Any optical brighteners or other brightening or whitening agents known in the art can be incorporated at levels typically from about 0.05% to about 1.2%, by weight, into the detergent compositions herein. Commercial optical brighteners which may be useful in the present invention can be classified into subgroups, which include, but are not necessarily limited to, derivatives of stilbene, pyrazoline, coumarin, carboxylic acid, methinecyanines, dibenzothiphene-5,5-dioxide, azoles, 5- and 6-membered-ring

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heterocycles, and other miscellaneous agents. Examples of such brighteners are disclosed in "The Production and Application of Fluorescent Brightening Agents", M. Zahradnik, Published by John Wiley & Sons, New York (1982).

Specific examples of optical brighteners which are useful in the present compositions are those identified in U.S. Patent 4,790,856, issued to Wixon on December 13, 1988. These brighteners include the PHORWHITE series of brighteners from Verona. Other brighteners disclosed in this reference include: Tinopal UNPA, Tinopal CBS and Tinopal 5BM; available from Ciba-Geigy; Artic White CC and Artic White CWD, available from Hilton-Davis, located in Italy; the 2-(4-stryl-phenyl)-2H-napthol[1,2-d]triazoles; 4,4'-bis- (1,2,3-triazol-2-yl)-stil- benes; 4,4'-bis(stryl)bisphenyls; and the aminocoumarins. Specific examples of these brighteners include 4-methyl-7-diethyl- amino coumarin; 1,2-bis(-venzimidazol-2-yl)ethylene; 1,3-diphenyl-phrazolines; 2,5-bis(benzoxazol-2-yl)thiophene; 2-stryl-napth-[1,2-d]oxazole; and 2-(stilbene-4-yl)-2H-naphtho- [1,2-d]triazole. See also U.S. Patent 3,646,015, issued February 29, 1972 to Hamilton. Anionic brighteners are preferred herein.

Suds Suppressors

Compounds for reducing or suppressing the formation of suds can be incorporated into the compositions of the present invention. Suds suppression can be of particular importance in the so-called "high concentration cleaning process" as described in U.S. 4,489,455 and 4,489,574 and in front-loading European-style washing machines.

A wide variety of materials may be used as suds suppressors, and suds suppressors are well known to those skilled in the art. See, for example, Kirk Othmer Encyclopedia of Chemical Technology, Third Edition, Volume 7, pages 430-447 (John Wiley & Sons, Inc., 1979). One category of suds suppressor of particular interest encompasses monocarboxylic fatty acid and soluble salts therein. See U.S. Patent 2,954,347, issued September 27, 1960 to Wayne St. John. The monocarboxylic fatty acids and salts thereof used as suds suppressor typically have hydrocarbyl chains of 10 to about 24 carbon atoms, preferably 12 to 18 carbon atoms. Suitable salts include the alkali metal salts such as sodium, potassium, and lithium salts, and ammonium and alkanolammonium salts.

The detergent compositions herein may also contain non-surfactant suds suppressors. These include, for example: high molecular weight hydrocarbons such as paraffin, fatty acid esters (e.g., fatty acid triglycerides), fatty acid esters of monovalent alcohols, aliphatic C₁₈-C₄₀ ketones (e.g., stearone), etc. Other suds inhibitors include N-alkylated amino triazines such as tri- to hexa-alkylmelamines or di- to tetra-alkyldiamine chlortriazines formed as products of cyanuric chloride with two or three

moles of a primary or secondary amine containing 1 to 24 carbon atoms, propylene oxide, and monostearyl phosphates such as monostearyl alcohol phosphate ester and monostearyl di-alkali metal (e.g., K, Na, and Li) phosphates and phosphate esters. The hydrocarbons such as paraffin and haloparaffin can be utilized in liquid form. The liquid hydrocarbons will be liquid at room temperature and atmospheric pressure, and will have a pour point in the range of about -40°C and about 50°C, and a minimum boiling point not less than about 110°C (atmospheric pressure). It is also known to utilize waxy hydrocarbons, preferably having a melting point below about 100°C. The hydrocarbons constitute a preferred category of suds suppressor for detergent compositions. Hydrocarbon suds suppressors are described, for example, in U.S. Patent 4,265,779, issued May 5, 1981 to Gandolfo et al. The hydrocarbons, thus, include aliphatic, alicyclic, aromatic, and heterocyclic saturated or unsaturated hydrocarbons having from about 12 to about 70 carbon atoms. The term "paraffin," as used in this suds suppressor discussion, is intended to include mixtures of true paraffins and cyclic hydrocarbons.

Another preferred category of non-surfactant suds suppressors comprises silicone suds suppressors. This category includes the use of polyorganosiloxane oils, such as polydimethylsiloxane, dispersions or emulsions of polyorganosiloxane oils or resins, and combinations of polyorganosiloxane with silica particles wherein the polyorganosiloxane is chemisorbed or fused onto the silica. Silicone suds suppressors are well known in the art and are, for example, disclosed in U.S. Patent 4,265,779, issued May 5, 1981 to Gandolfo et al and European Patent Application No. 89307851.9, published February 7, 1990, by Starch, M. S.

Other silicone suds suppressors are disclosed in U.S. Patent 3,455,839 which relates to compositions and processes for defoaming aqueous solutions by incorporating therein small amounts of polydimethylsiloxane fluids.

Mixtures of silicone and silanated silica are described, for instance, in German Patent Application DOS 2,124,526. Silicone defoamers and suds controlling agents in granular detergent compositions are disclosed in U.S. Patent 3,933,672, Bartolotta et al, and in U.S. Patent 4,652,392, Baginski et al, issued March 24, 1987.

An exemplary silicone based suds suppressor for use herein is a suds suppressing amount of a suds controlling agent consisting essentially of:

- (i) polydimethylsiloxane fluid having a viscosity of from about 20 cs. to about 1.500 cs. at 25°C:
- (ii) from about 5 to about 50 parts per 100 parts by weight of (i) of siloxane resin composed of (CH₃)₃SiO_{1/2} units of SiO₂ units in a ratio of from

(CH₃)₃ SiO_{1/2} units and to SiO₂ units of from about 0.6:1 to about 1.2:1; and

(iii) from about 1 to about 20 parts per 100 parts by weight of (i) of a solid silica gel.

In the preferred silicone suds suppressor used herein, the solvent for a continuous phase is made up of certain polyethylene glycols or polyethylene-polypropylene glycol copolymers or mixtures thereof (preferred), or polypropylene glycol. The primary silicone suds suppressor is branched/crosslinked and preferably not linear.

To illustrate this point further, typical liquid laundry detergent compositions with controlled suds will optionally comprise from about 0.001 to about 1, preferably from about 0.01 to about 0.7, most preferably from about 0.05 to about 0.5, weight % of said silicone suds suppressor, which comprises (1) a nonaqueous emulsion of a primary antifoam agent which is a mixture of (a) a polyorganosiloxane, (b) a resinous siloxane or a silicone resin-producing silicone compound, (c) a finely divided filler material, and (d) a catalyst to promote the reaction of mixture components (a), (b) and (c), to form silanolates; (2) at least one nonionic silicone surfactant; and (3) polyethylene glycol or a copolymer of polyethylene-polypropylene glycol having a solubility in water at room temperature of more than about 2 weight %; and without polypropylene glycol. Similar amounts can be used in granular compositions, gels, etc. See also U.S. Patents 4,978,471, Starch, issued December 18, 1990, and 4,983,316, Starch, issued January 8, 1991, 5,288,431, Huber et al., issued February 22, 1994, and U.S. Patents 4,639,489 and 4,749,740, Aizawa et al at column 1, line 46 through column 4, line 35.

The silicone suds suppressor herein preferably comprises polyethylene glycol and a copolymer of polyethylene glycol/polypropylene glycol, all having an average molecular weight of less than about 1,000, preferably between about 100 and 800. The polyethylene glycol and polyethylene/polypropylene copolymers herein have a solubility in water at room temperature of more than about 2 weight %, preferably more than about 5 weight %.

The preferred solvent herein is polyethylene glycol having an average molecular weight of less than about 1,000, more preferably between about 100 and 800, most preferably between 200 and 400, and a copolymer of polyethylene glycol/polypropylene glycol, preferably PPG 200/PEG 300. Preferred is a weight ratio of between about 1:1 and 1:10, most preferably between 1:3 and 1:6, of polyethylene glycol:copolymer of polyethylene-polypropylene glycol.

The preferred silicone suds suppressors used herein do not contain polypropylene glycol, particularly of 4,000 molecular weight. They also preferably do not contain block copolymers of ethylene oxide and propylene oxide, like PLURONIC L101.

Other suds suppressors useful herein comprise the secondary alcohols (e.g., 2-alkyl alkanols) and mixtures of such alcohols with silicone oils, such as the silicones disclosed in U.S. 4,798,679, 4,075,118 and EP 150,872. The secondary alcohols include the C₆-C₁₆ alkyl alcohols having a C₁-C₁₆ chain. A preferred alcohol is 2-butyl octanol, which is available from Condea under the trademark ISOFOL 12. Mixtures of secondary alcohols are available under the trademark ISALCHEM 123 from Enichem. Mixed suds suppressors typically comprise mixtures of alcohol + silicone at a weight ratio of 1:5 to 5:1.

For any detergent compositions to be used in automatic laundry washing machines, suds should not form to the extent that they overflow the washing machine. Suds suppressors, when utilized, are preferably present in a "suds suppressing amount. By "suds suppressing amount" is meant that the formulator of the composition can select an amount of this suds controlling agent that will sufficiently control the suds to result in a low-sudsing laundry detergent for use in automatic laundry washing machines.

The compositions herein will generally comprise from 0% to about 5% of suds suppressor. When utilized as suds suppressors, monocarboxylic fatty acids, and salts therein, will be present typically in amounts up to about 5%, by weight, of the detergent composition. Preferably, from about 0.5% to about 3% of fatty monocarboxylate suds suppressor is utilized. Silicone suds suppressors are typically utilized in amounts up to about 2.0%, by weight, of the detergent composition, although higher amounts may be used. This upper limit is practical in nature, due primarily to concern with keeping costs minimized and effectiveness of lower amounts for effectively controlling sudsing. Preferably from about 0.01% to about 1% of silicone suds suppressor is used, more preferably from about 0.25% to about 0.5%. As used herein, these weight percentage values include any silica that may be utilized in combination with polyorganosiloxane, as well as any adjunct materials that may be utilized. Monostearyl phosphate suds suppressors are generally utilized in amounts ranging from about 0.1% to about 2%, by weight, of the composition. Hydrocarbon suds suppressors are typically utilized in amounts ranging from about 0.01% to about 5.0%, although higher levels can be used. The alcohol suds suppressors are typically used at 0.2%-3% by weight of the finished compositions.

Fabric Softeners

Various through-the-wash fabric softeners, especially the impalpable smectite clays of U.S. Patent 4,062,647, Storm and Nirschl, issued December 13, 1977, as well as other softener clays known in the art, can optionally be used typically at levels of from about 0.5% to about 10% by weight in the present compositions to provide fabric softener benefits concurrently with fabric cleaning. Clay softeners can be used in combination with amine and cationic softeners as disclosed, for example, in U.S. Patent 4,375,416, Crisp et al, March 1, 1983 and U.S. Patent 4,291,071, Harris et al, issued September 22, 1981.

Dye Transfer Inhibiting Agents

The compositions of the present invention may also include one or more materials effective for inhibiting the transfer of dyes from one fabric to another during the cleaning process. Generally, such dye transfer inhibiting agents include polyvinyl pyrrolidone polymers, polyamine N-oxide polymers, copolymers of N-vinylpyrrolidone and N-vinylimidazole, manganese phthalocyanine, peroxidases, and mixtures thereof. If used, these agents typically comprise from about 0.01% to about 10% by weight of the composition, preferably from about 0.01% to about 5%, and more preferably from about 0.05% to about 2%.

More specifically, the polyamine N-oxide polymers preferred for use herein contain units having the following structural formula: R-A_X-P; wherein P is a polymerizable unit to which an N-O group can be attached or the N-O group can form part of the polymerizable unit or the N-O group can be attached to both units; A is one of the following structures: -NC(O)-, -C(O)O-, -S-, -O-, -N=; x is 0 or 1; and R is aliphatic, ethoxylated aliphatics, aromatics, heterocyclic or alicyclic groups or any combination thereof to which the nitrogen of the N-O group can be attached or the N-O group is part of these groups. Preferred polyamine N-oxides are those wherein R is a heterocyclic group such as pyridine, pyrrole, imidazole, pyrrolidine, piperidine and derivatives thereof.

The N-O group can be represented by the following general structures:

$$(R_1)_x - N - (R_2)_y;$$
 $= N - (R_1)_x$
 $(R_3)_z$

wherein R₁, R₂, R₃ are aliphatic, aromatic, heterocyclic or alicyclic groups or combinations thereof; x, y and z are 0 or 1; and the nitrogen of the N-O group can be attached or form part of any of the aforementioned groups. The amine oxide unit of the polyamine N-oxides has a pKa <10, preferably pKa <7, more preferred pKa <6.

Any polymer backbone can be used as long as the amine oxide polymer formed is water-soluble and has dye transfer inhibiting properties. Examples of suitable polymeric backbones are polyvinyls, polyalkylenes, polyesters, polyethers, polyamide, polyimides, polyacrylates and mixtures thereof. These polymers include random or block copolymers where one monomer type is an amine N-oxide and the other monomer type is an N-oxide. The amine N-oxide polymers typically have a ratio of amine to the amine N-oxide of 10:1 to 1:1,000,000. However, the number of amine oxide groups present in the polyamine oxide polymer can be varied by appropriate copolymerization or by an appropriate degree of N-oxidation. The polyamine oxides can be obtained in almost any degree of polymerization. Typically, the average molecular weight is within the range of 500 to 1,000,000; more preferred 1,000 to 500,000; most preferred 5,000 to 100,000. This preferred class of materials can be referred to as "PVNO".

The most preferred polyamine N-oxide useful in the detergent compositions herein is poly(4-vinylpyridine-N-oxide) which as an average molecular weight of about 50,000 and an amine to amine N-oxide ratio of about 1:4.

Copolymers of N-vinylpyrrolidone and N-vinylimidazole polymers (referred to as a class as "PVPVI") are also preferred for use herein. Preferably the PVPVI has an average molecular weight range from 5,000 to 1,000,000, more preferably from 5,000 to 200,000, and most preferably from 10,000 to 20,000. (The average molecular weight range is determined by light scattering as described in Barth, et al., Chemical Analysis, Vol 113. "Modern Methods of Polymer Characterization", the disclosures of which are incorporated herein by reference.) The PVPVI copolymers typically have a molar ratio of N-vinylimidazole to N-vinylpyrrolidone from 1:1 to 0.2:1, more preferably from 0.8:1 to 0.3:1, most preferably from 0.6:1 to 0.4:1. These copolymers can be either linear or branched.

The present invention compositions also may employ a polyvinylpyrrolidone ("PVP") having an average molecular weight of from about 5,000 to about 400,000, preferably from about 5,000 to about 200,000, and more preferably from about 5,000 to about 50,000. PVP's are known to persons skilled in the detergent field; see, for example, EP-A-262,897 and EP-A-256,696, incorporated herein by reference. Compositions containing PVP can also contain polyethylene glycol ("PEG") having an average molecular weight from about 500 to about 100,000, preferably from about 1,000 to about 10,000. Preferably, the ratio of PEG to PVP on a ppm basis delivered in wash solutions is from about 2:1 to about 50:1, and more preferably from about 3:1 to about 10:1.

The detergent compositions herein may also optionally contain from about 0.005% to 5% by weight of certain types of hydrophilic optical brighteners which also provide a dye transfer inhibition action. If used, the compositions herein will preferably comprise from about 0.01% to 1% by weight of such optical brighteners.

The hydrophilic optical brighteners useful in the present invention are those having the structural formula:

wherein R₁ is selected from anilino, N-2-bis-hydroxyethyl and NH-2-hydroxyethyl; R₂ is selected from N-2-bis-hydroxyethyl, N-2-hydroxyethyl-N-methylamino, morphilino, chloro and amino; and M is a salt-forming cation such as sodium or potassium.

When in the above formula, R₁ is anilino, R₂ is N-2-bis-hydroxyethyl and M is a cation such as sodium, the brightener is 4,4',-bis[(4-anilino-6-(N-2-bis-hydroxyethyl)-s-triazine-2-yl)amino]-2,2'-stilbenedisulfonic acid and disodium salt. This particular brightener species is commercially marketed under the tradename Tinopal-UNPA-GX by Ciba-Geigy Corporation. Tinopal-UNPA-GX is the preferred hydrophilic optical brightener useful in the detergent compositions herein.

When in the above formula, R₁ is anilino, R₂ is N-2-hydroxyethyl-N-2-methylamino and M is a cation such as sodium, the brightener is 4,4'-bis[(4-anilino-6-(N-2-hydroxyethyl-N-methylamino)-s-triazine-2-yl)amino]2,2'-stilbenedisulfonic acid disodium salt. This particular brightener species is commercially marketed under the tradename Tinopal 5BM-GX by Ciba-Geigy Corporation.

When in the above formula, R₁ is anilino, R₂ is morphilino and M is a cation such as sodium, the brightener is 4,4'-bis[(4-anilino-6-morphilino-s-triazine-2-yl)amino]2,2'-stilbenedisulfonic acid, sodium salt. This particular brightener species is commercially marketed under the tradename Tinopal AMS-GX by Ciba Geigy Corporation.

The specific optical brightener species selected for use in the present invention provide especially effective dye transfer inhibition performance benefits when used in combination with the selected polymeric dye transfer inhibiting agents hereinbefore described. The combination of such selected polymeric materials (e.g., PVNO and/or PVPVI) with such selected optical brighteners (e.g., Tinopal UNPA-GX, Tinopal 5BM-GX and/or Tinopal AMS-GX) provides significantly better dye transfer inhibition in

aqueous wash solutions than does either of these two detergent composition components when used alone. Without being bound by theory, it is believed that such brighteners work this way because they have high affinity for fabrics in the wash solution and therefore deposit relatively quick on these fabrics. The extent to which brighteners deposit on fabrics in the wash solution can be defined by a parameter called the "exhaustion coefficient". The exhaustion coefficient is in general as the ratio of a) the brightener material deposited on fabric to b) the initial brightener concentration in the wash liquor. Brighteners with relatively high exhaustion coefficients are the most suitable for inhibiting dye transfer in the context of the present invention.

Of course, it will be appreciated that other, conventional optical brightener types of compounds can optionally be used in the present compositions to provide conventional fabric "brightness" benefits, rather than a true dye transfer inhibiting effect. Such usage is conventional and well-known to detergent formulations.

Preparation of Cotton Soil Release Polymers EXAMPLE 1 PEI 1800 E7

The ethoxylation is conducted in a 2 gallon stirred stainless steel autoclave equipped for temperature measurement and control, pressure measurement, vacuum and inert gas purging, sampling, and for introduction of ethylene oxide as a liquid. A ~20 lb. net cylinder of ethylene oxide (ARC) is set up to deliver ethylene oxide as a liquid by a pump to the autoclave with the cylinder placed on a scale so that the weight change of the cylinder could be monitored.

A 750 g portion of polyethyleneimine (PEI) (Nippon Shokubai, Epomin SP-018 having a listed average molecular weight of 1800 equating to about 0.417 moles of polymer and 17.4 moles of nitrogen functions) is added to the autoclave. The autoclave is then sealed and purged of air (by applying vacuum to minus 28" Hg followed by pressurization with nitrogen to 250 psia, then venting to atmospheric pressure). The autoclave contents are heated to 130 °C while applying vacuum. After about one hour, the autoclave is charged with nitrogen to about 250 psia while cooling the autoclave to about 105 °C. Ethylene oxide is then added to the autoclave incrementally over time while closely monitoring the autoclave pressure, temperature, and ethylene oxide flow rate. The ethylene oxide pump is turned off and cooling is applied to limit any temperature increase resulting from any reaction exotherm. The temperature is maintained between 100 and 110 °C while the total pressure is allowed to gradually increase during the course of the reaction. After a total of 750 grams of ethylene oxide has been charged to the autoclave (roughly equivalent to one mole ethylene oxide per

PEI nitrogen function), the temperature is increased to 110 °C and the autoclave is allowed to stir for an additional hour. At this point, vacuum is applied to remove any residual unreacted ethylene oxide.

Next, vacuum is continuously applied while the autoclave is cooled to about 50 °C while introducing 376 g of a 25% sodium methoxide in methanol solution (1.74 moles, to achieve a 10% catalyst loading based upon PEI nitrogen functions). The methoxide solution is sucked into the autoclave under vacuum and then the autoclave temperature controller setpoint is increased to 130 °C. A device is used to monitor the power consumed by the agitator. The agitator power is monitored along with the temperature and pressure. Agitator power and temperature values gradually increase as methanol is removed from the autoclave and the viscosity of the mixture increases and stabilizes in about 1 hour indicating that most of the methanol has been removed. The mixture is further heated and agitated under vacuum for an additional 30 minutes.

Vacuum is removed and the autoclave is cooled to 105 °C while it is being charged with nitrogen to 250 psia and then vented to ambient pressure. The autoclave is charged to 200 psia with nitrogen. Ethylene oxide is again added to the autoclave incrementally as before while closely monitoring the autoclave pressure, temperature, and ethylene oxide flow rate while maintaining the temperature between 100 and 110 °C and limiting any temperature increases due to reaction exotherm. After the addition of 4500 g of ethylene oxide (resulting in a total of 7 moles of ethylene oxide per mole of PEI nitrogen function) is achieved over several hours, the temperature is increased to 110 °C and the mixture stirred for an additional hour.

The reaction mixture is then collected in nitrogen purged containers and eventually transferred into a 22 L three neck round bottomed flask equipped with heating and agitation. The strong alkali catalyst is neutralized by adding 167 g methanesulfonic acid (1.74 moles). The reaction mixture is then deodorized by passing about 100 cu. ft. of inert gas (argon or nitrogen) through a gas dispersion frit and through the reaction mixture while agitating and heating the mixture to 130 °C.

The final reaction product is cooled slightly and collected in glass containers purged with nitrogen.

In other preparations the neutralization and deodorization is accomplished in the reactor before discharging the product.

EXAMPLE 2

Quaternization of PEI 1800 E7

To a 500 mL Erlenmeyer flask equipped with a magnetic stirring bar is added polyethyleneimine having a molecular weight of 1800 which is further modified by

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ethoxylation to a degree of approximately 7 ethyleneoxy residues per nitrogen (PEI 1800, E7) (207.3g, 0.590 mol nitrogen, prepared as in Example I) and acetonitrile (120 g). Dimethyl sulfate (28.3g, 0.224 mol) is added in one portion to the rapidly stirring solution, which is then stoppered and stirred at room temperature overnight. The acetonitrile is removed by rotary evaporation at about 60°C, followed by further stripping of solvent using a Kugelrohr apparatus at approximately 80°C to afford 220 g of the desired partially quaternized material as a dark brown viscous liquid. The 13C-NMR (D₂O) spectrum obtained on a sample of the reaction product indicates the absence of a carbon resonance at ~58ppm corresponding to dimethyl sulfate. The ¹H-NMR (D₂O) spectrum shows a partial shifting of the resonance at about 2.5 ppm for methylenes adjacent to unquaternized nitrogen has shifted to approximately 3.0 ppm. This is consistent with the desired quaternization of about 38% of the nitrogens. **EXAMPLE 3**

Formation of amine oxide of PEI 1800 E7

To a 500 mL Erlenmeyer flask equipped with a magnetic stirring bar is added polyethyleneimine having a molecular weight of 1800 and ethoxylated to a degree of about 7 ethoxy groups per nitrogen (PEI-1800, E7) (209 g, 0.595 mol nitrogen, prepared as in Example I), and hydrogen peroxide (120 g of a 30 wt % solution in water, 1.06 mol). The flask is stoppered, and after an initial exotherm the solution is stirred at room temperature overnight. ¹H-NMR (D₂O) spectrum obtained on a sample of the reaction mixture indicates complete conversion. The resonances ascribed to methylene protons adjacent to unoxidized nitrogens have shifted from the original position at ~2.5 ppm to ~3.5 ppm. To the reaction solution is added approximately 5 g of 0.5% Pd on alumina pellets, and the solution is allowed to stand at room temperature for approximately 3 days. The solution is tested and found to be negative for peroxide by indicator paper. The material as obtained is suitably stored as a 51.1% active solution in water.

In other preparations, a lesser excess of hydrogen peroxide is used and the excess is left in the product or optionally destroyed by addition of a reducing agent such as sodium sulfite.

EXAMPLE 4 Formation of amine oxide of quaternized PEI 1800 E7

To a 500 mL Erlenmeyer flask equipped with a magnetic stirring bar is added polyethyleneimine having a molecular weight of 1800 which is further modified by ethoxylation to a degree of about 7 ethyleneoxy residues per nitrogen (PEI 1800 E7) and then further modified by quaternization to approximately 38% with dimethyl sulfate (130 g, ~0.20 mol oxidizeable nitrogen, hydrogen peroxide (48 g of a 30 wt % solution in water, 0.423 mol), and water (~50 g). The flask is stoppered, and after an initial

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exotherm the solution is stirred at room temperature overnight. ¹H-NMR (D₂O) spectrum obtained on a sample taken from the reaction mixture indicates complete conversion of the resonances attributed to the methylene peaks previously observed in the range of 2.5-3.0 ppm to a material having methylenes with a chemical shift of approximately 3.7 ppm. To the reaction solution is added approximately 5 g of 0.5% Pd on alumina pellets, and the solution is allowed to stand at room temperature for approximately 3 days. The solution is tested and found to be negative for peroxide by indicator paper. The desired material with ~38% of the nitrogens quaternized and 62% of the nitrogens oxidized to amine oxide is obtained and is suitably stored as a 44.9% active solution in water.

EXAMPLE 5

Preparation of PEI 1200 E7

The ethoxylation is conducted in a 2 gallon stirred stainless steel autoclave equipped for temperature measurement and control, pressure measurement, vacuum and inert gas purging, sampling, and for introduction of ethylene oxide as a liquid. A ~20 lb. net cylinder of ethylene oxide (ARC) is set up to deliver ethylene oxide as a liquid by a pump to the autoclave with the cylinder placed on a scale so that the weight change of the cylinder could be monitored.

A 750 g portion of polyethyleneimine (PEI) (having a listed average molecular weight of 1200 equating to about 0.625 moles of polymer and 17.4 moles of nitrogen functions) is added to the autoclave. The autoclave is then sealed and purged of air (by applying vacuum to minus 28" Hg followed by pressurization with nitrogen to 250 psia, then venting to atmospheric pressure). The autoclave contents are heated to 130 °C while applying vacuum. After about one hour, the autoclave is charged with nitrogen to about 250 psia while cooling the autoclave to about 105 °C. Ethylene oxide is then added to the autoclave incrementally over time while closely monitoring the autoclave pressure, temperature, and ethylene oxide flow rate. The ethylene oxide pump is turned off and cooling is applied to limit any temperature increase resulting from any reaction exotherm. The temperature is maintained between 100 and 110 °C while the total pressure is allowed to gradually increase during the course of the reaction. After a total of 750 grams of ethylene oxide has been charged to the autoclave (roughly equivalent to one mole ethylene oxide per PEI nitrogen function), the temperature is increased to 110 ° C and the autoclave is allowed to stir for an additional hour. At this point, vacuum is applied to remove any residual unreacted ethylene oxide.

Next, vacuum is continuously applied while the autoclave is cooled to about 50 ° C while introducing 376 g of a 25% sodium methoxide in methanol solution (1.74 moles, to achieve a 10% catalyst loading based upon PEI nitrogen functions). The methoxide solution is sucked into the autoclave under vacuum and then the autoclave temperature controller setpoint is increased to 130 °C. A device is used to monitor the power consumed by the agitator. The agitator power is monitored along with the temperature and pressure. Agitator power and temperature values gradually increase as methanol is removed from the autoclave and the viscosity of the mixture increases and stabilizes in about 1 hour indicating that most of the methanol has been removed. The mixture is further heated and agitated under vacuum for an additional 30 minutes.

Vacuum is removed and the autoclave is cooled to 105 °C while it is being charged with nitrogen to 250 psia and then vented to ambient pressure. The autoclave is charged to 200 psia with nitrogen. Ethylene oxide is again added to the autoclave incrementally as before while closely monitoring the autoclave pressure, temperature, and ethylene oxide flow rate while maintaining the temperature between 100 and 110 °C and limiting any temperature increases due to reaction exotherm. After the addition of 4500 g of ethylene oxide (resulting in a total of 7 moles of ethylene oxide per mole of PEI nitrogen function) is achieved over several hours, the temperature is increased to 110 °C and the mixture stirred for an additional hour.

The reaction mixture is then collected in nitrogen purged containers and eventually transferred into a 22 L three neck round bottomed flask equipped with heating and agitation. The strong alkali catalyst is neutralized by adding 167 g methanesulfonic acid (1.74 moles). The reaction mixture is then deodorized by passing about 100 cu. ft. of inert gas (argon or nitrogen) through a gas dispersion frit and through the reaction mixture while agitating and heating the mixture to 130 °C.

The final reaction product is cooled slightly and collected in glass containers purged with nitrogen.

In other preparations the neutralization and deodorization is accomplished in the reactor before discharging the product.

Other preferred examples such as PEI 1200 E15 and PEI 1200 E20 can be prepared by the above method by adjusting the reaction time and the relative amount of ethylene oxide used in the reaction.

EXAMPLE 6

9.7% Quaternization of PEI 1200 E7

To a 500ml erlenmeyer flask equipped with a magnetic stirring bar is added poly(ethyleneimine), MW 1200 ethoxylated to a degree of 7 (248.4g, 0.707 mol nitrogen, prepared as in Example 5) and acetonitrile (Baker, 200 mL). Dimethyl sulfate (Aldrich, 8.48g, 0.067 mol) is added all at once to the rapidly stirring solution, which is then

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stoppered and stirred at room temperature overnight. The acetonitrile is evaporated on the rotary evaporator at ~60°C, followed by a Kugelrohr apparatus (Aldrich) at ~80°C to afford ~220g of the desired material as a dark brown viscous liquid. A ¹³C-NMR (D₂O) spectrum shows the absence of a peak at ~58ppm corresponding to dimethyl sulfate. A ¹H-NMR (D₂O) spectrum shows the partial shifting of the peak at 2.5ppm (methylenes attached to unquaternized nitrogens) to ~3.0ppm.

EXAMPLE 7

Preparation of PEI 600 E₂₀

The ethoxylation is conducted in a 2 gallon stirred stainless steel autoclave equipped for temperature measurement and control, pressure measurement, vacuum and inert gas purging, sampling, and for introduction of ethylene oxide as a liquid. A \sim 20 lb. net cylinder of ethylene oxide (ARC) is set up to deliver ethylene oxide as a liquid by a pump to the autoclave with the cylinder placed on a scale so that the weight change of the cylinder could be monitored.

A 250 g portion of polyethyleneimine (PEI) (Nippon Shokubai, having a listed average molecular weight of 600 equating to about 0.417 moles of polymer and 6.25 moles of nitrogen functions) is added to the autoclave. The autoclave is then sealed and purged of air (by applying vacuum to minus 28" Hg followed by pressurization with nitrogen to 250 psia, then venting to atmospheric pressure). The autoclave contents are heated to 130 °C while applying vacuum. After about one hour, the autoclave is charged with nitrogen to about 250 psia while cooling the autoclave to about 105 °C. Ethylene oxide is then added to the autoclave incrementally over time while closely monitoring the autoclave pressure, temperature, and ethylene oxide flow rate. The ethylene oxide pump is turned off and cooling is applied to limit any temperature increase resulting from any reaction exotherm. The temperature is maintained between 100 and 110 °C while the total pressure is allowed to gradually increase during the course of the reaction. After a total of 275 grams of ethylene oxide has been charged to the autoclave (roughly equivalent to one mole ethylene oxide per PEI nitrogen function), the temperature is increased to 110 °C and the autoclave is allowed to stir for an additional hour. At this point, vacuum is applied to remove any residual unreacted ethylene oxide.

Next, vacuum is continuously applied while the autoclave is cooled to about 50 ° C while introducing 135 g of a 25% sodium methoxide in methanol solution (0.625 moles, to achieve a 10% catalyst loading based upon PEI nitrogen functions). The methoxide solution is sucked into the autoclave under vacuum and then the autoclave temperature controller setpoint is increased to 130 °C. A device is used to monitor the power consumed by the agitator. The agitator power is monitored along with the

temperature and pressure. Agitator power and temperature values gradually increase as methanol is removed from the autoclave and the viscosity of the mixture increases and stabilizes in about 1 hour indicating that most of the methanol has been removed. The mixture is further heated and agitated under vacuum for an additional 30 minutes.

Vacuum is removed and the autoclave is cooled to 105 °C while it is being charged with nitrogen to 250 psia and then vented to ambient pressure. The autoclave is charged to 200 psia with nitrogen. Ethylene oxide is again added to the autoclave incrementally as before while closely monitoring the autoclave pressure, temperature, and ethylene oxide flow rate while maintaining the temperature between 100 and 110 °C and limiting any temperature increases due to reaction exotherm. After the addition of approximately 5225 g of ethylene oxide (resulting in a total of 20 moles of ethylene oxide per mole of PEI nitrogen function) is achieved over several hours, the temperature is increased to 110 °C and the mixture stirred for an additional hour.

The reaction mixture is then collected in nitrogen purged containers and eventually transferred into a 22 L three neck round bottomed flask equipped with heating and agitation. The strong alkali catalyst is neutralized by adding 60 g methanesulfonic acid (0.625 moles). The reaction mixture is then deodorized by passing about 100 cu. ft. of inert gas (argon or nitrogen) through a gas dispersion frit and through the reaction mixture while agitating and heating the mixture to 130 °C.

The final reaction product is cooled slightly and collected in glass containers purged with nitrogen.

In other preparations the neutralization and deodorization is accomplished in the reactor before discharging the product.

<u>Preparation of Non-cotton Soil Release Polymers</u> EXAMPLE 8

Synthesis of Sodium 2-(2,3-Dihydroxypropoxy)ethanesulfonate Monomer To a 500ml, three neck, round bottom flask equipped with a magnetic stirring bar, modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-WatchTM, I²R) is added isethionic acid, sodium salt (Aldrich, 50.0g, 0.338 mol), sodium hydroxide (2.7g, 0.0675 mol), and glycerin (Baker, 310.9g, 3.38 mol). The solution is heated at 190°C under argon overnight as water distills from the reaction mixture. A ¹³C-NMR(DMSO-d₆) shows that the reaction is complete by the virtual disappearance of the isethionate peaks at ~53.5 ppm and ~57.4 ppm, and the emergence of product peaks at ~51.4 ppm (-CH₂SO₃Na) and ~67.5 ppm (CH₂CH₂SO₃Na). The solution is cooled to ~100°C and neutralized to pH 7 with methanesulfonic acid (Aldrich). The desired, neat material is obtained by adding 0.8

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mol% of potassium phosphate, monobasic as buffer and heating on a Kugelrohr apparatus (Aldrich) at 200°C for ~ 3 hrs. at ~1 mm Hg to afford 77g of yellow waxy solid. As an alternative, not all of the glycerin is removed before use in making the oligomers. The use of glycerin solutions of SEG can be a convenient way of handling this sulfonated monomer.

EXAMPLE 9

Synthesis of Sodium 2-[2-(2-Hydroxyethoxy)ethoxy]ethanesulfonate Monomer

To a 1L, three neck, round bottom flask equipped with a magnetic stirring bar, modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-Watch™, I2R) is added isethionic acid, sodium salt (Aldrich, 100.0g, 0.675 mol) and distilled water (~90 ml). After dissolution, one drop of hydrogen peroxide (Aldrich, 30% by wt. in water) is added to oxidize traces of bisulfite. The solution is stirred for one hour. A peroxide indicator strip shows a very weak positive test. Sodium hydroxide pellets (MCB, 2.5g, 0.0625 mol) are added, followed by diethylene glycol (Fisher, 303.3g, 2.86 mol). The solution is heated at 190C° under argon overnight as water distills from the reaction mixture. A 13C-NMR(DMSO-d6) shows that the reaction is complete by the disappearance of the isethionate peaks at ~53.5 ppm and ~57.4 ppm. The solution is cooled to room temperature and neutralized to pH 7 with 57.4g of a 16.4% solution of p-toluenesulfonic acid monohydrate in diethylene glycol. (Alternatively, methanesulfonic acid may be used.) The ¹³C-NMR spectrum of the product shows resonances at ~51ppm (-CH2SO3Na), ~60ppm (-CH2OH), and at ~69 ppm, ~72 ppm, and ~77 ppm for the remaining four methylenes. Small resonances are also visible for the sodium p-toluenesulfonate which formed during neutralization. The reaction affords 451g of a 35.3% solution of sodium 2-[2-(2hydroxyethoxy)ethoxylethanesulfonate in diethylene glycol. The excess diethylene glycol is removed by adding 0.8 mol% of monobasic potassium phosphate (Aldrich) as a buffer and heating on a Kugelrohr apparatus (Aldrich) at 150°C for ~ 3 hrs. at ~1 mm Hg to give the desired "SE3" (as defined herein above) as an extremely viscous oil or glass.

EXAMPLE 10

Synthesis of Sodium 2-{2-[2-(2-Hydroxyethoxy)ethoxy]ethoxy}ethoxy}ethoxy}ethoxy

To a 1L, three neck, round bottom flask equipped with a magnetic stirring bar, modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-Watch™, I²R) is added isethionic acid, sodium salt (Aldrich, 205.0g, 1.38 mol) and distilled water (~200 ml). After dissolution, one drop of hydrogen peroxide (Aldrich, 30% by wt. in water) is added to oxidize traces of bisulfite. The solution is stirred for one hour. A peroxide indicator strip shows a very weak positive

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test. Sodium hydroxide pellets (MCB, 5.5g, 0.138 mol) are added, followed by triethylene glycol (Aldrich, 448.7g, 3.0 mol). Optionally, the triethylene glycol can be purified by heating with strong base such as NaOH until color stabilizes and then distilling off the purified glycol for use in the synthesis. The solution is heated at 190°C under argon overnight as water distills from the reaction mixture. A 13C-NMR(DMSOd6) shows that the reaction is complete by the disappearance of the isethionate peaks at ~53.5 ppm and ~57.4 ppm, and the emergence of product peaks at ~51ppm (-CH₂SO₃Na), ~60ppm (-CH₂OH), and at ~67 ppm, ~69 ppm, and ~72 ppm for the remaining methylenes. The solution is cooled to room temperature and neutralized to pH 7 with methanesulfonic acid (Aldrich). The reaction affords 650g of a 59.5% solution of sodium 2-{2-[2-(2-hydroxyethoxy)ethoxy}ethoxy}ethoxy}ethoxy triethylene glycol. The excess triethylene glycol is removed by adding 0.8 mol% of monobasic potassium phosphate (Aldrich) as a buffer and heating on a Kugelrohr apparatus (Aldrich) at 180°C for ~ 5.5 hrs. at ~1 mm Hg to give the desired material as a brown solid. It is found that a more soluble buffer can be more effective in controlling pH during the stripping of excess triethylene glycol. One example of such a more soluble buffer is the salt of N-methylmorpholine with methanesulfonic acid. Alternatively, the pH can be controlled by frequent or continuous addition of acid such as methanesulfonic acid to maintain a pH near neutral during the stripping of excess glycol.

The material is believed to contain a low level of the disulfonate arising from reaction of both ends of the triethylene glycol with isethionate. However, the crude material is used without further purification as an anionic capping groups for polymer preparations.

Other preparations use a larger excess of triethylene glycol such as 5 to 10 moles per mole of isethionate.

EXAMPLE 11

Synthesis of an Oligomer of Sodium 2-[2-(2-Hydroxyethoxy)ethoxy]ethanesulfonate,

Dimethyl Terephthalate, Sodium 2-(2,3-Dihydroxypropoxy)ethanesulfonate, Glycerin,

Ethylene Glycol, and Propylene Glycol)

To a 250ml, three neck, round bottom flask equipped with a magnetic stirring bar, modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-Watch[®], I²R) is added sodium 2-[2-(2-hydroxyethoxy)ethoxy]ethanesulfonate (7.0g, 0.030 mol), dimethyl terephthalate (14.4g, 0.074 mol), sodium 2-(2,3-dihydroxypropoxy)ethanesulfonate (3.3g, 0.015 mol), glycerin (Baker, 1.4g, 0.015 mol), ethylene glycol (Baker, 14.0g, 0.225 mol), propylene

glycol (Fisher, 17.5g, 0.230 mol), and titanium (IV) propoxide (0.01g, 0.02% of total reaction weight). This mixture is heated to 180°C and maintained at that temperature overnight under argon as methanol and water distill from the reaction vessel. The material is transferred to a 500ml, single neck, round bottom flask and heated gradually over about 20 minutes to 240°C in a Kugelrohr apparatus (Aldrich) at about 2 mm Hg and maintained there for 1.5 hours. The reaction flask is then allowed to air cool quite rapidly to near room temperature under vacuum (~30 min.) The reaction affords 21.3g of the desired oligomer as a brown glass. A 13C-NMR(DMSO-d6) shows a resonance for -C(O)OCH2CH2O(O)C- at ~63.2 ppm (diester) and a resonance for -C(O)OCH2CH2OH at ~59.4 ppm (monoester). The ratio of the diester peak height to the monoester peak height is about 10. Resonances at ~51.5 ppm and ~51.6 ppm representing the sulfoethoxy groups (-CH2SO3Na) are also present. A ¹H-NMR(DMSO-d₆) shows a resonance at ~7.9 ppm representing terephthalate aromatic hydrogens. Analysis by hydrolysis-gas chromatography shows that the mole ratio of incorporated ethylene glycol to incorporated propylene glycol is 1.7:1. It also shows that about 0.9% of the final polymer weight consists of glycerin. If all glycerin monomer has been incorporated as esters of glycerin, it would represent approximately 4% of final oligomer weight. The solubility is tested by weighing a small amount of material into a vial, adding enough distilled water to make a 35% by weight solution, and agitating the vial vigorously. The material is readily soluble under these conditions.

EXAMPLE 12

Synthesis of an Oligomer of Sodium 2-[2-(2-Hydroxyethoxy)ethoxy]ethanesulfonate, Dimethyl Terephthalate, Sodium 2-(2,3-Dihydroxypropoxy)ethanesulfonate, Ethylene Glycol, and Propylene Glycol)

To a 250ml, three neck, round bottom flask equipped with a magnetic stirring bar, modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-Watch[®], I²R) is added sodium 2-[2-(2-hydroxyethoxy)ethoxy]ethanesulfonate (7.0g, 0.030 mol), dimethyl terephthalate (14.4g, 0.074 mol), sodium 2-(2,3-dihydroxypropoxy)ethanesulfonate (6.6g, 0.030 mol), ethylene glycol (Baker, 14.0g, 0.225 mol), propylene glycol (Fisher, 18.3g, 0.240 mol), and titanium (IV) propoxide (0.01g, 0.02% of total reaction weight). This mixture is heated to 180°C and maintained at that temperature overnight under argon as methanol distills from the reaction vessel. The material is transferred to a 500ml, single neck, round bottom flask and heated gradually over about 20 minutes to 240°C in a Kugelrohr apparatus (Aldrich) at about 0.1 mm Hg and maintained there for 110 minutes. The reaction flask is then allowed to air cool quite rapidly to near room temperature under

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vacuum (~30 min.) The reaction affords 24.4g of the desired oligomer as a brown glass. A 13C-NMR(DMSO-d₆) shows a resonance for -C(O)OCH2CH2O(O)C- at ~63.2 ppm (diester) and a resonance for -C(O)OCH2CH2OH at ~59.4 ppm (monoester). The ratio of the diester peak to monoester peak is measured to be 8. Resonances at ~51.5 ppm and ~51.6 ppm representing the sulfoethoxy groups (-CH2SO3Na) are also present. A ¹H-NMR(DMSO-d₆) shows a resonance at ~7.9 ppm representing terephthalate aromatic hydrogens. Analysis by Hydrolysis-GC shows that the mole ratio of incorporated ethylene glycol to incorporated propylene glycol is 1.6:1. The solubility is tested by weighing a small amount of material into a vial, adding enough distilled water to make a 35% by weight solution, and agitating the vial vigorously. The material is readily soluble under these conditions.

EXAMPLE 13

Synthesis of an Oligomer of Sodium 2-[2-(2-Hydroxyethoxy)ethoxy]ethanesulfonate, Dimethyl Terephthalate, Sodium 2-(2,3-Dihydroxypropoxy)ethanesulfonate, Glycerin, Ethylene Glycol, and Propylene Glycol)

To a 250ml, three neck, round bottom flask equipped with a magnetic stirring bar, modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-Watch®, I²R) is added sodium 2-[2-(2hydroxyethoxy)ethoxy]ethanesulfonate (7.0g, 0.030 mol), dimethyl terephthalate (9.6g, 0.049 mol), sodium 2-(2,3-dihydroxypropoxy)ethanesulfonate (2.2g, 0.010 mol), glycerin (Baker, 1.8g, 0.020 mol), ethylene glycol (Baker, 6.1g, 0.100 mol), propylene glycol (Fisher, 7.5g, 0.100 mol), and titanium (IV) propoxide (0.01g, 0.02% of total reaction weight). This mixture is heated to 180°C and maintained at that temperature overnight under argon as methanol distills from the reaction vessel. The material is transferred to a 250ml, single neck, round bottom flask and heated gradually over about 20 minutes to 240°C in a Kugelrohr apparatus (Aldrich) at about 3 mm Hg and maintained there for 1.5 hours. The reaction flask is then allowed to air cool quite rapidly to near room temperature under vacuum (~30 min.) The reaction affords 18.1g of the desired oligomer as a brown glass. A 13C-NMR(DMSO-d6) shows a resonance for -C(O)OCH2CH2O(O)C- at ~63.2 ppm (diester). A resonance for -C(O)OCH2CH2OH at ~59.4 ppm (monoester) is not detectable and is at least 12 times smaller than the diester peak. Resonances at ~51.5 ppm and ~51.6 ppm representing the sulfoethoxy groups (-CH2SO3Na) are also present. A 1H-NMR(DMSO-d6) shows a resonance at ~7.9 ppm representing terephthalate aromatic hydrogens. Analysis by Hydrolysis-GC shows that the mole ratio of incorporated ethylene glycol to incorporated propylene glycol is 1.6:1. The incorporated glycerin is found to be 0.45 weight% of the

final polymer. The solubility is tested by weighing a small amount of material into a vial, adding enough distilled water to make a 35% by weight solution, and agitating the vial vigorously. The material is readily soluble under these conditions.

EXAMPLE 14

Synthesis of an Oligomer of Sodium 2-[2-(2-Hydroxyethoxy)ethoxy]ethanesulfonate, Dimethyl Terephthalate, Sodium 2-(2,3-Dihydroxypropoxy)ethanesulfonate, Glycerol, Ethylene Glycol, and Propylene Glycol)

To a 250ml, three neck, round bottom flask equipped with a magnetic stirring bar. modified Claisen head, condenser (set for distillation), thermometer, and temperature controller (Therm-O-Watch®, I²R) is added sodium 2-I2-(2hydroxyethoxy)ethoxy]ethanesulfonate (2.7g, 0.011 mol, as in Example 2), dimethyl terephthalate (12.0g, 0.062 mol, Aldrich), sodium 2-(2,3dihydroxypropoxy)ethanesulfonate (5.0g, 0.022 mol, as in Example 1), glycerol (Baker, 0.50g, 0.0055 mol), ethylene glycol (Baker, 6.8g, 0.110 mol), propylene glycol (Baker, 8.5g, 0.112 mol), and titanium (IV) propoxide (0.01g, 0.02% of total reaction weight). This mixture is heated to 180°C and maintained at that temperature overnight under argon as methanol and water distill from the reaction vessel. The material is transferred to a 500ml, single neck, round bottom flask and heated gradually over about 20 minutes to 240°C in a Kugelrohr apparatus (Aldrich) at about 0.5 mm Hg and maintained there for 150 minutes. The reaction flask is then allowed to air cool quite rapidly to near room temperature under vacuum (~30 min.) The reaction affords 16.7g of the desired oligomer as a brown glass. A 13C-NMR(DMSO-d6) shows a resonance for -C(O)OCH2CH2O(O)C- at ~63.2 ppm (diester) and a resonance for -C(O)OCH2CH2OH at ~59.4 ppm (monoester). The ratio of the peak height for the diester resonance to that of the monoester resonance is measured to be 6.1. Resonances at ~51.5 ppm and ~51.6 ppm representing the sulfoethoxy groups (-CH2SO3Na) are also present. A ¹H-NMR(DMSO-d₆) shows a resonance at ~7.9 ppm representing terephthalate aromatic hydrogens. Analysis by hydrolysis-gas chromatography shows that the mole ratio of incorporated ethylene glycol to incorporated propylene glycol is 1.42:1. The solubility is tested by weighing a small amount of material into a vial, adding enough distilled water to make a 35% by weight solution, and agitating the vial vigorously. The material is readily soluble under these conditions. A ~9g sample of this material is further heated at 240°C in a Kugelrohr apparatus at about 0.5 mm Hg and maintained there for 80 minutes. A ¹³C-NMR(DMSO-d₆) shows no detectable peak for monoester at ~59.4 ppm. The peak for diester at ~63.2 ppm is at least 11 times larger than the monoester

peak. The solubility of this material is tested as above and it is also found to be readily soluble under these conditions.

The following describe high density liquid detergent compositions according to the present invention:

TABLE I

	weight %			
Ingredient	15	16	17	18
Polyhydroxy Coco-Fatty Acid Amide	3.65	3.50		
C ₁₂ -C ₁₃ Alcohol Ethoxylate E ₉	3.65	0.80	••	
Sodium C ₁₂ -C ₁₅ Alcohol Sulfate	6.03	2.50		
Sodium C ₁₂ -C ₁₅ Alcohol Ethoxylate	9.29	15.10		
E _{2.5} Sulfate				
Sodium C ₁₄ -C ₁₅ Alcohol Ethoxylate			18.00	18.00
E _{2.25} Sulfate				
Alkyl N-Methyl Glucose Amide			4.50	4.50
C ₁₀ Amidopropyl Amine		1.30		••
Citric Acid	2.44	3.00	3.00	3.00
Fatty Acid (C ₁₂ -C ₁₄)	4.23	2.00	2.00	2.00
NEODOL 23-91			2.00	2.00
Ethanol	3.00	2.81	3.40	3.40
Monoethanolamine	1.50	0.75	1.00	1.00
Propanediol	8.00	7.50	7.50	7.00
Boric Acid	3.50	3.50	3.50	3.50
Tetraethylenepentamine		1.18		
Sodium Toluene Sulfonate	2.50	2.25	2.50	2.50
NaOH	2.08	2.43	2.62	2.62
Minors ²	1.60	1.30	0.27	0.27
Cotton Soil Release Polymer ³	0.50	0.50		
Cotton Soil Release Polymer ⁴		-	2.00	1.00
Non-cotton Soil Release Polymer ⁵	0.33	0.22	·	••
Non-cotton Soil Release Polymer ⁶			1.00	
Non-cotton Soil Release Polymer ⁷			••	1.00
Water ⁸	balance	balance	balance	balance

^{1.} E9 Ethoxylated Alcohols as sold by the Shell Oil Co.

- 2. Minors includes optical brightener and enzymes (protease, lipase, cellulase, and amylase).
- 3. Cotton soil release polymer according to Example 7.
- 4. Cotton soil release polymer according to Example 1.
- 5. Non-cotton soil release polymer according to U.S. Patent 4,968,451, Scheibel et al., issued November 6, 1990.
- 6. Non-cotton soil release polymer according to Example 11.
- 7. Non-cotton soil release polymer according to U.S. Patent 4,702,857, Gosselink, issued October 27, 1987.
- 8. Balance to 100% can, for example, include minors like optical brightener, perfume, suds suppresser, soil dispersant, protease, lipase, cellulase, chelating agents, dye transfer inhibiting agents, additional water, and fillers, including CaCO₃, talc, silicates, etc.

TABLE II

	weight %			
<u>Ingredient</u>	19	20	21	22
Polyhydroxy Coco-Fatty Acid Amide	2.50	2.50	4->	**
C ₁₂ -C ₁₃ Alcohol Ethoxylate E9		==	3.65	0.80
Sodium C ₁₂ -C ₁₅ Alcohol Sulfate			6.03	2.50
Sodium C ₁₂ -C ₁₅ Alcohol Ethoxylate	20.15	20.15	-	
E _{1.8} Sulfate				
Sodium C ₁₄ -C ₁₅ Alcohol Ethoxylate			18.00	18.00
E _{2.25} Sulfate				
Alkyl N-Methyl Glucose Amide			4.50	4.50
C ₁₀ Amidopropyl Amine	0.50	0.50		
Citric Acid	2.44	3.00	3.00	3.00
Fatty Acid (C ₁₂ -C ₁₄)			2.00	2.00
NEODOL 23-91	0.63	0.63		
Ethanol	3.00	2.81	3.40	3.40
Monoethanolamine	1.50	0.75	1.00	1.00
Propanediol	8.00	7.50	7.50	7.00
Boric Acid	3.50	3.50	3.50	3.50
Ethoxylate tetraethylenepentamine ²	0.50		-	0.50
Tetraethylenepentamine		1.18		
Sodium Toluene Sulfonate	2.50	2.25	2.50	2.50

NaOH	2.08	2.43	2.62	2.62
Minors ³	1.60	1.30	0.27	0.27
Cotton Soil Release Polymer ⁴	0.50	0.50		
Cotton Soil Release Polymer ⁵			2.00	1.00
Non-cotton Soil Release Polymer ⁶	0.33			0.55
Non-cotton Soil Release Polymer ⁷		0.50	1.00	
Non-cotton Soil Release Polymer ⁸	••			1.00
Water ⁹	balance	balance	balance	balance

- 1. Eq Ethoxylated Alcohols as sold by the Shell Oil Co.
- 2. Ethoxylated tetraethylenepentamine (PEI 189 E₁₅-E₁₈) according to U. S. 4,597,898 Vander Meer issued July 1, 1986.
- 3. Minors includes optical brightener and enzymes (protease, lipase, cellulase, and amylase).
- 4. Cotton soil release polymer according to Example 7.
- 5. Cotton soil release polymer according to Example 4.
- 6. Non-cotton soil release polymer according to U.S. Patent 4,968,451, Scheibel et al., issued November 6, 1990.
- 7. Non-cotton soil release polymer according to Example 11.
- 8. Non-cotton soil release polymer according to U.S. Patent 4,702,857, Gosselink, issued October 27, 1987.
- 9. Balance to 100% can, for example, include minors like optical brightener, perfume, suds suppresser, soil dispersant, protease, lipase, cellulase, chelating agents, dye transfer inhibiting agents, additional water, and fillers, including CaCO₃, talc, silicates, etc.

TABLE III

Ingredients	23	24	25	26
Sodium C ₁₄ -C ₁₅ Alcohol Ethoxylate	13.00	••	••	8.43
E _{2.25} Sulfate				
Sodium C ₁₂ -C ₁₅ Alcohol Ethoxylate		18.00	13.00	
E _{2.5} Sulfate				
Sodium C ₁₂ -C ₁₃ linear alkylbenzene	9.86			8.43
sulfonate				
Fatty Acid (C ₁₂ -C ₁₄)		2.00	2.00	2.95
C ₁₂ -C ₁₃ Alcohol Ethoxylate E ₉				3.37
C ₁₀ Amidopropyl Amine			0.80	

NEODOL 23-91	2.22	2.00	1.60	
Alkyl N-Methyl Glucose Amide		5.00	2.50	
Citric Acid	7.10	3.00	3.00	3.37
Ethanol	1.92	3.52	3.41	1.47
Monoethanolamine	0.71	1.09	1.00	1.05
Propanediol	4.86	8.00	6.51	6.00
Boric Acid	2.22	3.30	2.50	
Ethoxylated Tetraethylenepentamine	1.18	1.18		1.48
Sodium Cumene Sulfonate	1.80	3.00		3.00
Sodium Toluene Sulfonate			2.50	
NaOH	6.60	2.82	2.90	2.10
Dodecyltrimethylammonium Chloride		_		0.51
Sodium Tartrate Mono and Di-succinate		***		3.37
Sodium Formate				0.32
Minors ²	1.60	1.80	2.00	1.60
Cotton Soil Release Polymer ³	0.50	2.00		
Cotton Soil Release Polymer ⁴	1.50		2.00	3.00
Non-cotton Soil Release Polymer ⁵	1.50		2.00	••
Non-cotton Soil Release Polymer ⁶		1.15		1.50
Water ⁷	balance	balance	balance	balance

- 1. Eq Ethoxylated Alcohols as sold by the Shell Oil Co.
- 2. Minors includes optical brightener and enzymes (protease, lipase, cellulase, and amylase).
- 3. Cotton soil release polymer according to Example 4.
- 4. Cotton soil release polymer according to Example 7.
- 5. Non-cotton soil release polymer according to Example 10.
- 6. Non-cotton soil release polymer according to Example 11.
- 7. Balance to 100% can, for example, include minors like optical brightener, perfume, suds suppresser, soil dispersant, protease, lipase, cellulase, chelating agents, dye transfer inhibiting agents, additional water, and fillers, including CaCO₃, talc, silicates, etc.

TABLE IV

Ingredients 27 28 29 30

	12.00			
Sodium C ₁₄ -C ₁₅ Alcohol Ethoxylate	13.00			8.43
E _{2.25} Sulfate				
Sodium C ₁₂ -C ₁₅ Alcohol Ethoxylate		18.00	13.00	
E _{2,5} Sulfate				
Sodium C ₁₂ -C ₁₃ linear alkylbenzene	9.86			8.43
sulfonate				
Fatty Acid (C ₁₂ -C ₁₄)		2.00	2.00	2.95
C ₁₂ -C ₁₃ Alcohol Ethoxylate E9			••	3.37
C ₁₀ Amidopropyl Amine			0.80	
NEODOL 23-9 ¹	2.22	2.00	1.60	
Alkyl N-Methyl Glucose Amide		5.00	2.50	
Citric Acid	7.10	3.00	3.00	3.37
Ethanol	1.92	3.52	3.41	1.47
Monoethanolamine	0.71	1.09	1.00	1.05
Propanediol	4.86	8.00	6.51	6.00
Boric Acid	2.22	3.30	2.50	
Ethoxylated Tetraethylenepentamine	1.18	1.18	ţ	1.48
Sodium Cumene Sulfonate	1.80	3.00		3.00
Sodium Toluene Sulfonate		••	2.50	
NaOH	6.60	2.82	2.90	2.10
Dodecyltrimethylammonium Chloride				0.51
Sodium Tartrate Mono and Di-succinate				3.37
Sodium Formate				0.32
Minors ²	1.60	1.80	2.00	1.60
Cotton Soil Release Polymer ³	0.50	2.00		••
Cotton Soil Release Polymer ⁴	1.50		2.00	3.00
Non-cotton Soil Release Polymer ⁵	1.50		2.00	
Non-cotton Soil Release Polymer ⁶		1.15		1.50
Water ⁷	balance	balance	balance	balance

- 1. E9 Ethoxylated Alcohols as sold by the Shell Oil Co.
- 2. Minors includes optical brightener and enzymes (protease, lipase, cellulase, and amylase).
- 3. Cotton soil release polymer according to Example 7.
- 4. Cotton soil release polymer according to Example 1.
- 5. Non-cotton soil release polymer according to Example 10.

- 6. Non-cotton soil release polymer according to Example 11.
- 7. Balance to 100% can, for example, include minors like optical brightener, perfume, suds suppresser, soil dispersant, protease, lipase, cellulase, chelating agents, dye transfer inhibiting agents, additional water, and fillers, including CaCO₃, talc, silicates, etc.

TABLE V

Ingredients	31	32	33	34	35
Polyhydroxy coco-fatty acid	3.50	3.50	3.15	3.50	3.00
amide					
NEODOL 23-9 ¹	2.00	0.60	2.00	0.60	0.60
C ₂₅ Alkyl ethoxylate sulphate	19.00	19.40	19.00	17.40	14.00
C ₂₅ Alkyl sulfate				2.85	2.30
C ₁₀ -Aminopropylamide				0.75	0.50
Citric acid	3.00	3.00	3.00	3.00	3.00
Tallow fatty acid	2.00	2.00	2.00	2.00	2.00
Ethanol	3.41	3.47	3.34	3.59	2.93
Propanediol	6.22	6.35	6.21	6.56	5.75
Monomethanol amine	1.00	0.50	0.50	0.50	0.50
Sodium hydroxide	3.05	2.40	2.40	2.40	2.40
Sodium p-toluene sulfonate	2.50	2.25	2.25	2.25	2.25
Borax	2.50	2.50	2.50	2.50	2.50
Protease 2	0.88	0.88	0.88	0.88	0.88
Lipolase ³	0.04	0.12	0.12	0.12	0.12
Duramyl ⁴	0.10	0.10	0.10	0.10	0.40
CAREZYME	0.053	0.053	0.053	0.053	0.053
Optical Brightener	0.15	0.15	0.15	0.15	0.15
Cotton soil release agent 5	1.18	1.18	1.18	1.18	1.75
Non-cotton soil release agent 6	0.22	0.15	0.15	0.15	0.15
Fumed silica	0.119	0.119	0.119	0.119	0.119
Minors, aestetics, water	balance	balance	balance	balance	balance

- 1. C_{12} - C_{13} alkyl E9 ethoxylate as sold by Shell Oil Co.
- 2. Bacillus amyloliquefaciens subtilisin as described in WO 95/10615 published April 20, 1995 by Genencor International.
- 3. Derived from Humicola lanuginosa and commercially available from Novo.

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- 4. Disclosed in WO 9510603 A and available from Novo.
- 5. PEI 1800 E7.
- 6. Terephthalate co-polymer as disclosed in U.S. Patent 4,968,451, Scheibel et al., issued November 6, 1990.

WHAT IS CLAIMED IS:

- 1. A liquid laundry detergent composition comprising:
 - A) at least 0.01% by weight, of an anionic detersive surfactant selected from the group consisting of alkyl sulfates, alkyl alkoxylated sulfates, and mixtures thereof;
 - B) at least 0.01% by weight, of a non-cotton soil release agent selected from the group consisting of a terephthalate co-polymer comprising:
 - i) a backbone comprising:
 - a) at least one moiety having the formula:

b) at least one moiety having the formula:

wherein R⁹ is C₂-C₆ linear alkylene, C₃-C₆ branched alkylene, C₅-C₇ cyclic alkylene, and mixtures thereof; R¹⁰ is independently selected from hydrogen or -L-SO₃-M⁺; wherein L is a side chain moiety selected from the group consisting of alkylene, oxyalkylene, alkyleneoxyalkylene, arylene, oxyarylene, alkyleneoxyarylene, poly(oxyalkylene), oxy-alkyleneoxyarylene, poly(oxyalkylene)oxyarlyene, alkylene-poly(oxyalkylene)oxyarlyene, alkylene-poly(oxyalkylene), and mixtures thereof; M is hydrogen or a salt forming cation; i has the value of 0 or 1;

- at least one trifunctional, ester-forming, branching moiety;
- d) at least one 1,2-oxyalkyleneoxy moiety; and
- ii) one or more capping units comprising:

1 3 1

- a) ethoxylated or propoxylated hydroxyethanesulfonate or ethoxylated or propoxylated hydroxypropanesulfonate units of the formula (MO₃S)(CH₂)_m(R¹¹O)_n-, where M is a salt forming cation, R¹¹ is ethylene, propylene, and mixtures thereof, m is 0 or 1, and n is from 1 to 20;
- b) sulfoaroyl units of the formula -(O)C(C₆H₄)(SO₃-M⁺), wherein M is a salt forming cation;
- c) modified poly(oxyethylene)oxy monoalkyl ether units of the formula R¹²O(CH₂CH₂O)_k-, wherein R¹² contains from 1 to 4 carbon atoms and k is from 3 to 100; and
- ethoxylated or propoxylated phenolsulfonate endcapping units of the formula MO3S(C6H4)(OR¹³)nO-, wherein n is from 1 to 20; M is a salt-forming cation; and R¹³ is ethylene, propylene, and mixtures thereof;

a sulfonated oligomeric ester composition comprising the sulfonated product of a pre-formed, substantially linear ester oligomer, said linear ester oligomer comprising, per mole,

- i) 2 moles of terminal units wherein from 1 mole to 2 moles of said terminal units are derived from an olefinically unsaturated component selected from the group consisting of allyl alcohol and methallyl alcohol, and any remaining of said terminal units are other units of said linear ester oligomer;
- ii) from 1 mole to 4 moles of nonionic hydrophile units, said hydrophile units being derived from alkyleneoxides, said alkylene oxides comprising from 50% to 100% ethylene oxide:
- iii) from 1.1 moles to 20 moles of repeat units derived from an aryldicarbonyl component wherein said aryldicarbonyl component is comprised of from 50% to 100% dimethylterephthalate, whereby the repeat units derived from said dimethylterephthalate are terephthaloyl; and
- iv) from 0.1 moles to 19 moles of repeat units derived from a diol component selected from the group consisting of C₂-C₄ glycols;

wherein the extent of sulfonation of said sulfonated oligomeric ester composition is such that said terminal units are chemically modified by

v) from 1 mole to 4 moles of terminal unit substituent groups of formula -SO_XM wherein x is 2 or 3, said terminal unit substituent groups being derived from a bisulfite component selected from the group consisting of HSO₃M wherein M is a conventional water-soluble cation;

a capped terephthalate co-polymer having the formula

$$X[(OCH_2CH_2)_n(OR_5)_m][(A-R^1-A-R^2)_{tt}(A-R^3-A-R^2)_v]$$

 $A-R^4-A[(R^5O)_m(CH_2CH_2O)_n]X$

wherein each of the A moieties is selected from the group consisting of

and combinations thereof, each of the R¹ moieties is selected from the group consisting of 1.4-phenylene and combinations thereof with 1,3-phenylene, 1,2 phenylene, 1,8-naphthylene, 1,4-naphthylene, 2,2'biphenylene, 4,4'-biphenylene, C1-C8 alkylene, C1-C8 alkenylene and mixtures thereof the R2 moieties are each selected from the group consisting of ethylene moieties, substituted ethylene moieties having C₁-C₄ alkyl, alkoxy substituents, and mixtures thereof; the R³ moieties are substituted C₂-C₁₈ hydrocarbylene moieties having A)]w[(R⁵O)_m(CH₂CH₂O)_n]X substituent; the R⁴ moieties are R¹ or R³ moieties, or mixtures thereof; each R⁵ is C₁-C₄ alkylene, or the moiety -R²-A-R⁶- wherein R⁶ is a C₁-C₁₂ alkylene, alkenylene, arylene, or alkarylene moiety; each M is hydrogen or a water-soluble cation; each X is C₁-C₄ alkyl; the indices m and n have the values such that the moiety -(CH2CH2O)- comprises at least 50% by weight of the moiety [(R⁵O)_m(CH₂CH₂O)_n], provided that when R⁵ is the moiety -R²-A-R⁶-, m is 1; each n is at least 10; the indices u and v have the value such that the sum of u + v is from 3 to 25; the index w is 0 or at least 1; and when w is at least 1 u, v and w have the value such that the sum of u + v + w is from 3 to 25; and mixtures thereof:

c) at least 0.01% by weight, of a water-soluble or dispersible, modified polyamine cotton soil release agent comprising a polyamine backbone corresponding to the formula:

$$H_{2N-R}$$
_{n+1}-[N-R]_m-[N-R]_n-NH₂

having a modified polyamine formula $V_{(n+1)}W_mY_nZ$ or a polyamine backbone corresponding to the formula:

having a modified polyamine formula $V_{(n-k+1)}W_mY_nY_kZ$, wherein k is less than or equal to n, said polyamine backbone prior to modification has a molecular weight greater than 200 daltons, wherein

i) V units are terminal units having the formula:

ii) W units are backbone units having the formula:

iii) Y units are branching units having the formula:

$$-N-R- \quad \text{or} \quad -N^+-R- \quad \text{or} \quad -N-R- \quad ; \text{and}$$

iv) Z units are terminal units having the formula:

wherein backbone linking R units are selected from the group consisting of C2-C12 alkylene, C4-C12 alkenylene, C3-C12 hydroxyalkylene, C₄-C₁₂ dihydroxy-alkylene, C₈-C₁₂ dialkylarylene, $-(R^{1}O)_{x}R^{1}$ -, $-(R^{1}O)_{x}R^{5}(OR^{1})_{x}$ -, - $(CH_2CH(OR^2)CH_2O)_z^{-}(R^1O)_vR^1(OCH_2CH(OR^2)CH_2)_{w^{-}}$, - $C(O)(R^4)_{r}C(O)_{r}$ -CH₂CH(OR²)CH₂-, and mixtures thereof, preferably C2-C12 alkylene, C3-C12 hydroxyalkylene, C4-C₁₂ dihydroxyalkylene, C₈-C₁₂ dialkylarylene, -(R¹O)_xR¹-, - $(R^{1}O)_{x}R^{5}(OR^{1})_{x}$ $(CH_2CH(OH)CH_2O)_z(R^1O)_vR^1(OCH_2CH(OH)CH_2)_{w^*}$ CH₂CH(OR²)CH₂-, and mixtures thereof, more preferably C₂-C₁₂ alkylene, C3-C12 hydroxyalkylene, C4-C12 dihydroxyalkylene, $-(R^{1}O)_{x}R^{1}$ -, $-(R^{1}O)_{x}R^{5}$ - $(OR^{1})_{x}$ -, $-(CH_{2}CH(OH)CH_{2}O)_{z}$ -(R¹O)_vR¹(OCH₂CH(OH)CH₂)_w, and mixtures thereof, most preferably C₂-C₁₂ alkylene, and mixtures thereof; wherein R¹ is C₂-C₆ alkylene, preferably ethylene, and mixtures thereof; R² is hydrogen, -(R¹O)_xB, and mixtures thereof, preferably hydrogen; R³ is C1-C18 alkyl, C7-C12 arylalkyl, C7-C12 alkyl substituted aryl, C₆-C₁₂ aryl, and mixtures thereof, C₁-C₆ alkyl and mixtures thereof, more preferably methyl; R⁴ is C₁-C₁₂ alkylene, C₄-C₁₂ alkenylene, C₈-C₁₂ arylalkylene, C₆-C₁₀ arylene, and mixtures thereof, preferably C2-C12 alkylene, C8-C12 arylalkylene, and mixtures thereof, more preferably, ethylene, butylene, and mixtures thereof; R⁵ is C₁-C₁₂ alkylene, C₃-C₁₂ hydroxyalkylene, C₄-C₁₂ dihydroxy-alkylene, C8-C12 dialkylarylene, -C(O)-, - $C(O)NHR^6NHC(O)$ -, $-R^1(OR^1)$ -, $-C(O)(R^4)$ _rC(O)-, -CH₂CH(OH)CH₂-, CH2CH(OH)CH2O(R1O)_VR1OCH2CH(OH)CH2-, and mixtures thereof, preferably ethylene, -C(O)-, -C(O)NHR⁶NHC(O)-, $R^{1}(OR^{1})_{v^{-}}$, -(CH₂CH(OH)CH₂O)_z($R^{1}O$)_y $R^{1}(OCH_{2}CH(OH)-$ CH₂)_w -, -CH₂CH(OH)CH₂-, and mixtures thereof, more preferably -CH₂CH(OH)CH₂-; R⁶ is C₂-C₁₂ alkylene or C₆-C₁₂ arylene; E units are selected from the group consisting of hydrogen, C₁-C₂₂

alkyl, C3-C22 alkenyl, C7-C22 arylalkyl, C2-C22 hydroxyalkyl, - $(CH_2)_pCO_2M$, - $(CH_2)_qSO_3M$, - $CH(CH_2CO_2M)CO_2M$, (CH₂)_pPO₃M, -(R¹O)_xB, -C(O)R³, and mixtures thereof; preferably hydrogen, C_3 - C_{22} hydroxyalkyl, benzyl, C_1 - C_{22} alkyl, - $(R^1O)_xB$, - $C(O)R^3$, $-(CH_2)_pCO_2^-M^+$, $-(CH_2)_qSO_3^-M^+$, $-CH(CH_2CO_2M)$ -CO₂M and mixtures thereof, more preferably hydrogen, C₁-C₂₂ alkyl, -(R¹O)_xB, -C(O)R³, and mixtures thereof, most preferably -(R¹O)_xB; provided that when any E unit of a nitrogen is a hydrogen, said nitrogen is not also an N-oxide; B is hydrogen, C1-C6 alkyl, - $(\mathsf{CH}_2)_q \mathsf{SO}_3 \mathsf{M}, -(\mathsf{CH}_2)_p - \mathsf{CO}_2 \mathsf{M}, -(\mathsf{CH}_2)_q (\mathsf{CHSO}_3 \mathsf{M}) \mathsf{CH}_2 \mathsf{SO}_3 \mathsf{M}, -(\mathsf{CH}_2)_q \mathsf{M}_3 \mathsf{M}_3$ (CH₂)_q(CHSO₂M)CH₂-SO₃M, -(CH₂)_pPO₃M, -PO₃M, and mixtures thereof, preferably hydrogen, C1-C6 alkyl, -(CH2)qSO3M, -(CH₂)_q(CHSO₃M)-CH₂SO₃M, -(CH₂)_q(CHSO₂M)CH₂SO₃M, and mixtures thereof, more preferably hydrogen, -(CH2)qSO3M, and mixtures thereof, most preferably hydrogen; M is hydrogen or a water soluble cation in sufficient amount to satisfy charge balance; X is a water soluble anion; m has the value from 4 to 400; n has the value from 0 to 200; p has the value from 1 to 6, q has the value from 0 to 6; r has the value of 0 or 1; w has the value 0 or 1; x has the value from 1 to 100; y has the value from 0 to 100; z has the value 0 or 1: and

- D) the balance carrier and adjunct ingredients wherein said composition has a pH of 7.2 to 8.9 when measured as a 10% solution in water.
- 2. A composition according to Claim 1 further comprising a nonionic surfactant, selected from the group consisting of alkyl alkoxylate, a fatty acid amide having the formula:

wherein R⁷ is C₇-C₂₂ alkyl, R⁸ is independently selected from the group consisting of hydrogen, C₁-C₄ alkyl, C₁-C₄ hydroxyalkyl, -(C₂H₄O)_jH, and mixtures thereof; wherein j is from 1 to 3; and mixtures of said surfactants, preferably alkyl alkoxylate, a fatty acid amide having the formula:

wherein R^7 is C_7 - C_{22} alkyl, R^8 is independently selected from the group consisting of hydrogen, C_1 - C_4 alkyl, C_1 - C_4 hydroxyalkyl, -(C_2 H₄O)_jH, and mixtures thereof; wherein j is from 1 to 3; and mixtures of said surfactants.

- 3. A composition according to either of Claims 1 or 2 wherein the non-cotton soil release polymer is a member selected from the group consisting of:
 - I) a non-cotton soil release agent comprising:
 - a) a backbone comprising:
 - i) at least one moiety having the formula:

ii) at least one moiety having the formula:

wherein R⁹ is C₂-C₆ linear alkylene, C₃-C₆ branched alkylene, C₅-C₇ cyclic alkylene, and mixtures thereof; R¹⁰ is independently selected from hydrogen or -L-SO₃-M⁺; wherein L is a side chain moiety selected from the group consisting of alkylene, oxyalkylene, alkyleneoxyalkylene, arylene, oxyarylene, alkyleneoxyarylene, poly(oxyalkylene), oxy-alkyleneoxyarylene, poly(oxyalkylene)oxyarlyene, alkylene-poly(oxyalkylene), and mixtures thereof; M is hydrogen or a salt forming cation; i has the value of 0 or 1;

- iii) at least one trifunctional, ester-forming, branching moiety;
- iv) at least one 1,2-oxyalkyleneoxy moiety; and
- b) one or more capping units comprising:
 - ethoxylated or propoxylated hydroxyethanesulfonate or ethoxylated or propoxylated

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hydroxypropanesulfonate units of the formula (MO₃S)(CH₂)_m(R¹¹O)_n-, where M is a salt forming cation, R¹¹ is ethylene, propylene, and mixtures thereof, m is 0 or 1, and n is from 1 to 20;

- sulfoaroyl units of the formula -(O)C(C6H4)(SO3ii) M⁺), wherein M is a salt forming cation;
- iii) modified poly(oxyethylene)oxy monoalkyl ether units of the formula R¹²O(CH₂CH₂O)_k-, wherein R¹² contains from 1 to 4 carbon atoms and k is from 3 to 100; and
- iv) ethoxylated or propoxylated phenolsulfonate endcapping units of the formula MO₃S(C₆H₄)-(OR¹³)_nO-, wherein n is from 1 to 20; M is a saltforming cation; and R¹³ is ethylene, propylene, and mixtures thereof;
- II) a sulfonated oligomeric ester composition comprising the sulfonated product of a pre-formed, substantially linear ester oligomer, said linear ester oligomer comprising, per mole,
 - 2 moles of terminal units wherein from 1 mole to 2 moles of a) said terminal units are derived from an olefinically unsaturated component selected from the group consisting of allyl alcohol and methallyl alcohol, and any remaining of said terminal units are other units of said linear ester oligomer;
 - from 1 mole to 4 moles of nonionic hydrophile units, said b) hydrophile units being derived from alkyleneoxides, said alkylene oxides comprising from 50% to 100% ethylene oxide;
 - from 1.1 moles to 20 moles of repeat units derived from an c) aryldicarbonyl component wherein said aryldicarbonyl component is comprised of from 50% to 100% dimethylterephthalate, whereby the repeat units derived from said dimethylterephthalate are terephthaloyl; and
 - from 0.1 moles to 19 moles of repeat units derived from a diol d) component selected from the group consisting of C2-C4 glycols:

wherein the extent of sulfonation of said sulfonated oligomeric ester composition is such that said terminal units are chemically modified by

- e) from 1 mole to 4 moles of terminal unit substituent groups of formula -SO_XM wherein x is 2 or 3, said terminal unit substituent groups being derived from a bisulfite component selected from the group consisting of HSO₃M wherein M is a conventional water-soluble cation; and
- III) a non-cotton soil release agent of the formula

$$X[(OCH_2CH_2)_n(OR_5)_m][(A-R^1-A-R^2)_u(A-R^3-A-R^2)_v]$$

$$A-R^4-A[(R^5O)_m(CH_2CH_2O)_m]X$$

wherein each of the A moieties is selected from the group consisting of

and combinations thereof, each of the R1 moieties is selected from the group consisting of 1,4-phenylene and combinations thereof with 1,3-phenylene, 1,2 phenylene, 1,8-naphthylene, 1,4-naphthylene, 2,2'biphenylene, 4,4'-biphenylene, C1-C8 alkylene, C1-C8 alkenylene and mixtures thereof the R² moieties are each selected from the group consisting of ethylene moieties, substituted ethylene moieties having C1-C4 alkyl, alkoxy substituents, and mixtures thereof; the R³ moieties are substituted C₂-C₁₈ hydrocarbylene moieties having at least one -CO₂M, -O[(R^5O)_m(CH₂CH₂O)_n]X or -A[(R^2 -A- R^4 -A)]w[(R⁵O)_m(CH₂CH₂O)_n]X substituent; the R⁴ moieties are R¹ or R³ moieties, or mixtures thereof; each R⁵ is C₁-C₄ alkylene, or the moiety -R²-A-R⁶- wherein R⁶ is a C₁-C₁₂ alkylene, alkenylene, arylene, or alkarylene moiety; each M is hydrogen or a water-soluble cation; each X is C₁-C₄ alkyl; the indices m and n have the values such that the moiety -(CH2CH2O)- comprises at least 50% by weight of the moiety [(R⁵O)_m(CH₂CH₂O)_n], provided that when R⁵ is the moiety -R²-A-R⁶-, m is 1; each n is at least 10; the indices u and v have the value such that the sum of u + v is from 3 to 25; the index w is 0 or at least 1; and when w is at least 1 u, v and w have the value

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such that the sum of u + v + w is from 3 to 25; and mixtures of said non-cotton soil release agents.

- 4. A composition according to any of Claims 1-3 wherein said adjunct ingredients are selected from the group consisting of builders, enzymes, enzyme stabilizers, optical brighteners, bleaches, bleach boosters, bleach activators, dye transfer agents, dispersents, enzyme activators, suds suppressers, dyes, perfumes, colorants, filler salts, hydrotropes, and mixtures thereof.
- 5. A liquid laundry detergent composition according to any of Claims 1-4 further comprising a protease enzyme.

INTERNATIONAL SEARCH REPORT

Intern 1al Application No PCT/US 97/06918

A. CLASS IPC 6	BIFICATION OF SUBJECT MATTER C11D3/37		
According	to International Patent Classification (IPC) or to both national class	ification and IPC	
B. FIELD	S SEARCHED		
	documentation searched (classification system followed by classific	pation symbols)	
IPC 6	C11D		
Document	ation searched other than minimum documentation to the extent the	at such documents are included in the fields sea	rched
Electronia	data base consulted during the international search (name of data	base and, where practical, search terms used)	
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT		
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A	US 4 891 160 A (VANDER MEER JAN January 1990 cited in the application see claims	1ES M) 2	1,4,5
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X Fun	rther documents are listed in the continuation of box O.	X Patent family members are listed	in annex.
* Special o	satisfaction of cited documents :	"T" later document published after the inter	
"A" doour	nent defining the general state of the art which is not idened to be of cartioular relevance	or priority date and not in conflict with cited to understand the principle or the	
'E' earlier	r document but published on or after the international	invention "X" document of particular relevance; the o	daimed invention
"L" doour	date nent which may throw doubts on priority claim(s) or	cannot be considered novel or cannot knows an inventive step when the do	t be considered to
	h is cited to establish the publication date of another on or other special receon (as specified)	"Y" document of particular relevance; the o	laimed invention
	ment referring to an oral disclosure, use, exhibition or r means	document is combined with one or mo ments, such combination being obvio	ore other such doou-
P docum	nent published prior to the international filing data but than the priority date claimed	in the art. "&" document member of the same patent	•
	ectual completion of the international search	Date of mailing of the international sea	
	26 September 1997	0 8, 10, 97	
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentiass 2	Authorized officer	
	ML - 2280 HV Rijawijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Grittern, A	

INTERNATIONAL SEARCH REPORT

Intern val Application No PCT/US 97/06918

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C.(Continu	ation) DOCUMENTS C NSIDERED TO BE RELEVANT	
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